

GenCore version 5.1.6  
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OM nucleic - nucleic search, using sw model

Run on: July 1, 2004, 18:34:00 ; Search time 5328 Seconds  
(without alignments)  
10746.274 Million cell updates/sec

Title: US-09-934-249-1  
Perfect score: 1321  
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Scoring table: COLIGO\_NUC  
Gapop 60.0 , Gapext 60.0

Searched: 3470272 seqs, 21671516995 residues

Word size : 0

Total number of hits satisfying chosen parameters: 1603530

Minimum DB seq length: 0  
Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database : GenEmbl.\*

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- 6: gb.pat.\*
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- 10: gb.ro.\*
- 11: gb.sts.\*
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- 15: em.ba.\*
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- 40: em.hugo.mus.\*
- 41: em.hugo.other.\*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed,  
and is derived by analysis of the total score distribution.

SUMMARIES

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| 3          | 24    | 1.8           | 50     | 6  | AR232180 | AR232180 Sequence  |
| 4          | 23    | 1.7           | 25     | 6  | AR028113 | AR028113 Sequence  |
| 5          | 23    | 1.7           | 25     | 6  | AR030289 | AR030289 Sequence  |
| 6          | 23    | 1.7           | 25     | 6  | I42108   | I42108 Sequence 3  |
| 7          | 23    | 1.7           | 30     | 6  | I84401   | I84401 Sequence 2  |
| 8          | 23    | 1.7           | 31     | 6  | A01419   | A01419 Malaria par |
| 9          | 23    | 1.7           | 33     | 6  | I84406   | I84406 Sequence 7  |
| 10         | 23    | 1.7           | 45     | 6  | AR366218 | AR366218 Sequence  |
| 11         | 23    | 1.7           | 47     | 6  | AX539586 | AX539586 Sequence  |
| 12         | 23    | 1.7           | 47     | 6  | AX539587 | AX539587 Sequence  |
| 13         | 23    | 1.7           | 48     | 6  | AR366215 | AR366215 Sequence  |
| 14         | 23    | 1.7           | 48     | 6  | AX539588 | AX539588 Sequence  |
| 15         | 23    | 1.7           | 48     | 6  | AX539589 | AX539589 Sequence  |
| 16         | 23    | 1.7           | 50     | 6  | AX539582 | AX539582 Sequence  |
| 17         | 23    | 1.7           | 50     | 6  | AX539583 | AX539583 Sequence  |
| 18         | 22    | 1.7           | 22     | 6  | AR336835 | AR336835 Sequence  |
| 19         | 22    | 1.7           | 38     | 6  | AX539590 | AX539590 Sequence  |
| 20         | 22    | 1.7           | 38     | 6  | AX539591 | AX539591 Sequence  |
| 21         | 22    | 1.7           | 39     | 6  | AX539592 | AX539592 Sequence  |
| 22         | 22    | 1.7           | 39     | 6  | AX539593 | AX539593 Sequence  |
| 23         | 21    | 1.6           | 21     | 6  | AR084563 | AR084563 Sequence  |
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| 25         | 21    | 1.6           | 21     | 6  | AR084567 | AR084567 Sequence  |
| 26         | 21    | 1.6           | 21     | 6  | AR084578 | AR084578 Sequence  |
| 27         | 21    | 1.6           | 21     | 6  | AR084579 | AR084579 Sequence  |
| 28         | 21    | 1.6           | 21     | 6  | AR084582 | AR084582 Sequence  |
| 29         | 21    | 1.6           | 21     | 6  | AR093142 | AR093142 Sequence  |
| 30         | 21    | 1.6           | 39     | 6  | E50506   | E50506 Method for  |
| 31         | 21    | 1.6           | 39     | 6  | E50507   | E50507 Method for  |
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| 36         | 20    | 1.5           | 20     | 6  | AX104051 | AX104051 Sequence  |
| 37         | 20    | 1.5           | 20     | 6  | AX355382 | AX355382 Sequence  |
| 38         | 20    | 1.5           | 20     | 6  | AX547104 | AX547104 Sequence  |
| 39         | 20    | 1.5           | 20     | 6  | BD069976 | BD069976 Use of nu |
| 40         | 20    | 1.5           | 24     | 6  | AR078306 | AR078306 Sequence  |
| 41         | 20    | 1.5           | 24     | 6  | AR078307 | AR078307 Sequence  |
| 42         | 20    | 1.5           | 28     | 6  | AX034217 | AX034217 Sequence  |
| 43         | 20    | 1.5           | 28     | 6  | A62658   | A62658 Sequence 22 |
| 44         | 20    | 1.5           | 50     | 6  | AR144730 | AR144730 Sequence  |
| 45         | 20    | 1.5           | 50     | 6  | AR232179 | AR232179 Sequence  |

ALIGNMENTS

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| RESULT 1   | AX199566   | AX199566    | 50 bp | DNA | linear | PAT 29-AUG-2001 |
| LOCUS      | Sequence 496 from Patent WO0151670.  |             |       |     |        |                 |
| DEFINITION | AX199566   |             |       |     |        |                 |
| ACCESSION  | AX199566.1   | GI:15389997 |       |     |        |                 |
| VERSION    |  |             |       |     |        |                 |
| KEYWORDS   | Homo sapiens (human)   |             |       |     |        |                 |
| SOURCE     | Homo sapiens   |             |       |     |        |                 |
| ORGANISM   | Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo. |             |       |     |        |                 |
| REFERENCE  | Shimkets, R.A. and Leach, M.D.   |             |       |     |        |                 |
| AUTHORS    | Nucleic acids containing single nucleotide polymorphisms and   |             |       |     |        |                 |
| TITLE      | methods of use thereof   |             |       |     |        |                 |

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JOURNAL Patent: WO 0151670-A 496 19-JUL-2001;
FEATURES Curagen Corporation (US)
source Location/Qualifiers
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/mol_type="unassigned DNA"
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misc_feature 25..26
/note="Nucleotide deleted between bases 25 and 26
Accession number cg42747251"
misc_feature 26
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Db 1 CACGGTCTTCATCAGCGGCACAGC 26
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LOCUS AR242044 30 bp DNA linear PAT 20-DEC-2002
DEFINITION Sequence 332 from patent US 6472154.
ACCESSION AR242044
VERSION AR242044.1 GI:27287856
KEYWORDS Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 30)
AUTHORS Garner,H.R., Wren,J.D., Minna,J.D. and Fondon,J.W. III.
TITLE Polymorphic repeats in human genes
JOURNAL Patent: US 6472154-A 332 29-OCT-2002;
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QY 56 GCGCGCGCGCGCGCGCGCGCGG 79
Db 7 GCGCGCGCGCGCGCGCGCGCGG 30
RESULT 3
LOCUS AR232180 50 bp DNA linear PAT 20-DEC-2002
DEFINITION Sequence 14 from patent US 6455292.
ACCESSION AR232180
VERSION AR232180.1 GI:27274071
KEYWORDS Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 50)
AUTHORS Shu,Y., Fan,W., Kovacs,K.P., Zidanic,M. and Jay,G.
TITLE Full-length serine protein kinase in brain and pancreas
JOURNAL Patent: US 6455292-A 14 24-SEP-2002;
FEATURES Location/Qualifiers
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LOCUS AR028113 25 bp DNA linear PAT 29-SEP-1999
DEFINITION Sequence 3 from patent US 5858649.
ACCESSION AR028113
VERSION AR028113.1 GI:5940086
KEYWORDS Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 25)
AUTHORS Asgari,M., Blick,M., Bresser,J., Cubbage,M.Lee. and Prashad,N.
TITLE Amplification of mRNA for distinguishing fetal cells in maternal blood
JOURNAL Patent: US 5858649-A 3 12-JAN-1999;
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ACCESSION AR030289
VERSION AR030289.1 GI:5943503
KEYWORDS Unknown.
ORGANISM Unknown.
REFERENCE 1 (bases 1 to 25)
AUTHORS Asgari,M., Blick,M., Bresser,J., Cubbage,M.Lee. and Prashad,N.
TITLE Intracellular antigens for identifying fetal cells in maternal blood
JOURNAL Patent: US 5861253-A 3 19-JAN-1999;
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LOCUS I42108 25 bp DNA linear PAT 07-OCT-1997
DEFINITION Sequence 3 from patent US 5629147.
ACCESSION I42108
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ACCESSION AX539586  
VERSION AX539586.1 GI:25273135  
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SOURCE synthetic construct  
ORGANISM synthetic construct  
artificial sequences.  
REFERENCE 1  
TITLE Brinkmann,U., Hoffmeyer,S. and Mornhinweg,E.  
Polymorphisms in the human gene for the multidrug  
resistance-associated protein 1 (mrp-1) and their use in diagnostic  
and therapeutic applications  
JOURNAL Patent: WO 02059142-A 373 01-AUG-2002;  
Epidaurus Biotechnologie AG (DE)  
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LOCUS AX539587 47 bp DNA linear PAT 23-NOV-2002  
DEFINITION Sequence 374 from Patent WO02059142.  
ACCESSION AX539587  
VERSION AX539587.1 GI:25273137  
KEYWORDS  
SOURCE synthetic construct  
ORGANISM synthetic construct  
artificial sequences.  
REFERENCE 1  
AUTHORS Brinkmann,U., Hoffmeyer,S. and Mornhinweg,E.  
TITLE Polymorphisms in the human gene for the multidrug  
resistance-associated protein 1 (mrp-1) and their use in diagnostic  
and therapeutic applications  
JOURNAL Patent: WO 02059142-A 374 01-AUG-2002;  
Epidaurus Biotechnologie AG (DE)  
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AR366215  
LOCUS AR366215 48 bp DNA linear PAT 12-SEP-2003  
DEFINITION Sequence 19 from patent US 6329147.  
ACCESSION AR366215  
VERSION AR366215.1 GI:34598597  
KEYWORDS  
SOURCE Unknown.  
ORGANISM Unknown.  
REFERENCE 1 (bases 1 to 48)  
Unclassified.

AUTHORS Wagner,R.E. Jr.  
TITLE Methods for detection of a triplet repeat block and a functional  
mismatch binding protein in a biological fluid sample  
JOURNAL Patent: US 6329147-A 19 11-DEC-2001;  
FEATURES  
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ACCESSION AX539588  
VERSION AX539588.1 GI:25273139  
KEYWORDS  
SOURCE synthetic construct  
ORGANISM synthetic construct  
artificial sequences.  
REFERENCE 1  
AUTHORS Brinkmann,U., Hoffmeyer,S. and Mornhinweg,E.  
TITLE Polymorphisms in the human gene for the multidrug  
resistance-associated protein 1 (mrp-1) and their use in diagnostic  
and therapeutic applications  
JOURNAL Patent: WO 02059142-A 375 01-AUG-2002;  
Epidaurus Biotechnologie AG (DE)  
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LOCUS AX539589 48 bp DNA linear PAT 23-NOV-2002  
DEFINITION Sequence 376 from Patent WO02059142.  
ACCESSION AX539589  
VERSION AX539589.1 GI:25273140  
KEYWORDS  
SOURCE synthetic construct  
ORGANISM synthetic construct  
artificial sequences.  
REFERENCE 1  
AUTHORS Brinkmann,U., Hoffmeyer,S. and Mornhinweg,E.  
TITLE Polymorphisms in the human gene for the multidrug  
resistance-associated protein 1 (mrp-1) and their use in diagnostic  
and therapeutic applications  
JOURNAL Patent: WO 02059142-A 376 01-AUG-2002;  
Epidaurus Biotechnologie AG (DE)  
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Search completed: July 1, 2004, 20:48:03  
Job time : 5330 secs

GenCore version 5.1.6  
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OM nucleic - nucleic search, using sw model

Run on: July 1, 2004, 18:02:49 ; Search time 580 Seconds  
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9675.637 Million cell updates/sec

Title: US-09-934-249-1  
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Scoring table: OLIGO NUC  
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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| 3          | 24    | 1.8         | 30     | 7     | Abx80007 EST polym |
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| 6          | 23    | 1.7         | 25     | 2     | Aaq55856 Fragile X |
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| 14         | 23    | 1.7         | 48     | 6     | Aad40337 Test DNA  |
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| 16         | 23    | 1.7         | 48     | 6     | Abz67038 Human MRP |
| 17         | 23    | 1.7         | 50     | 6     | Abz67031 Human MRP |
| 18         | 23    | 1.7         | 50     | 6     | Abz67032 Human MRP |
| 19         | 23    | 1.7         | 50     | 7     | Abz22157 E. coli i |
| 20         | 22    | 1.7         | 22     | 9     | Aad60109 Human PME |
| 21         | 22    | 1.7         | 38     | 6     | Abz67040 Human MRP |
| 22         | 22    | 1.7         | 38     | 6     | Abz67039 Human MRP |
| 23         | 22    | 1.7         | 39     | 6     | Abz67042 Human MRP |

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| C | 25 | 22 | 1.7 | 41 | 6 | ABZ47846 | Abz47846 Human ATP |
|   | 26 | 22 | 1.7 | 41 | 6 | AAK99304 | AAK99304 Human can |
|   | 27 | 22 | 1.7 | 41 | 6 | AAK99303 | AAK99303 Human can |
| C | 28 | 21 | 1.6 | 21 | 3 | AZ44349  | Az44349 Protein k  |
| C | 29 | 21 | 1.6 | 21 | 6 | ABK99279 | ABK99279 Hepatitis |
| C | 30 | 21 | 1.6 | 21 | 9 | ADC16572 | Adc16572 Short int |
|   | 31 | 21 | 1.6 | 21 | 9 | ADC16571 | Adc16571 Short int |
| C | 32 | 21 | 1.6 | 39 | 4 | AAH27630 | Aah27630 Upper str |
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|   | 34 | 20 | 1.5 | 20 | 2 | AAV47686 | Aav47686 Umethyla  |
|   | 35 | 20 | 1.5 | 20 | 2 | AAV74243 | Aav74243 CpG-N mot |
|   | 36 | 20 | 1.5 | 20 | 4 | AAF99116 | Aaf99116 Immunosti |
|   | 37 | 20 | 1.5 | 20 | 6 | ABS77759 | Abz77759 Angiogene |
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|   | 45 | 20 | 1.5 | 33 | 2 | AAQ73441 | Aaq73441 Crohn's d |

## ALIGNMENTS

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AC AAH89715;  
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DT 01-OCT-2001 (first entry)  
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DE Human coding sequence polymorphic site SEQ ID NO: 496.  
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KW Human; single nucleotide polymorphism; SNP; paternity test;  
forensic test; aberrant protein expression; ds.  
XX  
OS Homo sapiens.  
XX  
FN WO200151670-A2.  
PD 19-JUL-2001.  
XX  
PF 05-JAN-2001; 2001WO-US000322.  
XX  
PR 07-JAN-2000; 2000US-0174962P.  
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PA (CURA-) CURAGEN CORP.  
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PI Shimkets RA, Leach MD;  
XX  
DR WPI; 2001-451871/48.  
XX  
PT P-PSDB; AAM00598.  
PT Isolated human polynucleotides containing single nucleotide  
polymorphisms, useful for the treatment and diagnosis of e.g. cancer,  
infection and diabetes.  
XX  
FS Claim 1; Page 246; 475pp; English.  
XX  
CC The present invention relates to human nucleic acids containing single  
nucleotide polymorphisms (SNPs). These can be used in forensic and  
paternity tests, and to aid in the treatment of diseases associated with  
aberrant protein expression, including cancer, amyloidosis, diabetes,  
Alzheimer's disease, Down's syndrome, oedema, lupus (SLE), vasculitis,  
glomerulonephritis, haemolytic anaemia, thrombocytopaenia, arthritis,  
meningitis, muscular disorders, dementia, neurological diseases, tubercous  
sclerosis, male infertility, hypercalcaemia, blood pressure disorders,  
osteoporosis, pathogenic infections, hypercholesterolaemia, obesity or  
autoimmunity. The present sequence is a polymorphism-containing

CC oligonucleotide fragment of the invention  
 XX  
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 RESULT 2  
 ABZ84399/c  
 ID ABZ84399 standard; DNA; 25 BP.  
 XX  
 AC ABZ84399;  
 XX  
 DT 14-MAY-2003 (first entry)  
 DE  
 XX Toxicologically relevant human PCR primer #1558.  
 DE  
 XX Toxicologically relevant gene; toxicological response; PCR primer; ss.  
 KW  
 XX Homo sapiens;  
 OS Synthetic.  
 OS  
 XX WO2003016500;A2.  
 PN  
 XX 27-FEB-2003.  
 PD  
 XX 16-AUG-2002; 2002WO-US026514.  
 PF  
 XX 16-AUG-2001; 2001US-0313080P.  
 PR  
 XX (PHAS-) PHASE-1 MOLECULAR TOXICOLOGY INC.  
 PA  
 XX Neft RE, Dunn RT, Adkins K, Pickett GG, Kier LD, Schweiser K;  
 PI Alen P;  
 PI  
 XX WPI; 2003-268322/26.  
 DR  
 XX  
 XX Determining a toxicological response to an agent, useful for screening of  
 PT drugs, comprises comparing the expression profile of one or more human  
 PT toxic response genes to a reference gene expression profile indicative of  
 PT toxicity.  
 PT  
 XX Claim 1; Page 346; 455pp; English.  
 PS  
 XX The present invention describes a method (M1) for determining a  
 CC toxicological response to an agent, which comprises comparing the  
 CC expression profile of one or more human toxic response genes to a  
 CC reference gene expression profile indicative of toxicity, and so  
 CC determining the presence of a toxic response to the agent. Also  
 CC described: (1) an array comprising one or more polynucleotides selected  
 CC from the genes corresponding to the partial sequences given in ABZ84399  
 CC to ABZ84764, or their fragments of at least 20 nucleotides, or homologues  
 CC; and (2) determining if a gene putatively identified to be a toxic  
 CC response gene plays a role on toxic response pathways by determining the  
 CC expression profile of the gene after exposure of cells or a human subject  
 CC to a known toxic pharmaceutical or industrial agent, comprising: (a)  
 CC exposing cells to an agent or isolating cells from a human subject who  
 CC was exposed to an agent; (b) obtaining the test gene expression profile  
 CC for a putatively identified toxic response gene after exposure to a known  
 CC toxic pharmaceutical or industrial agent; and (c) comparing the test  
 CC profile to the expression profile of a gene with a similar function or  
 CC comparing the test profile to the expression profile of that gene after  
 CC exposure to other known toxic compounds. The methods are useful for  
 CC predicting and determining toxicological responses on a cellular, organ  
 CC or system level. The arrays comprising the human genes are useful for  
 CC toxicological screening of drugs, pharmaceutical compounds and chemicals

SQ Sequence 25 BP; 4 A; 6 C; 9 G; 6 T; 0 U; 0 Other;  
 Query Match 1.9%; Score 25; DB 7; Length 25;  
 Best Local Similarity 100.0%; Pred. No. 0.62;  
 Matches 25; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 1105 CTACAGCGAGGTCTCGGCCACTAC 1129  
 DB 25 CTACAGCGAGGTCTCGGCCACTAC 1  
 RESULT 3  
 ABX80007  
 ID ABX80007 standard; cDNA; 30 BP.  
 XX  
 AC ABX80007;  
 XX  
 DT 17-APR-2003 (first entry)  
 DE  
 XX EST polymorphic DNA repeat polynucleotide #332.  
 DE  
 XX EST; expressed sequence tag; ss; polymorphic repeat; tandem repeat;  
 KW polymorphic marker prediction of ubiquitous simple sequences; POMPOUS;  
 KW Rep-X; human; genetic disease; drug-treatment; Machado-Joseph;  
 KW Haw River syndrome; Huntington's disease; fragile-X syndrome;  
 KW Predreich's ataxia; myotonic dystrophy; hyperandrogenaemia;  
 KW spinal atrophy; bulbar atrophy; spinocerebellar ataxia.  
 KW  
 XX Homo sapiens.  
 OS  
 XX US6472154-B1.  
 PN  
 XX 29-OCT-2002.  
 PD  
 XX 31-DEC-1999; 99US-00475947.  
 PF  
 XX 31-DEC-1999; 99US-00475947.  
 PR  
 XX (TEXA) UNIV TEXAS SYSTEM.  
 PA  
 XX Garner HR, Wren JD, Minna JD, Fondon JW;  
 PI  
 XX WPI; 2003-208818/20.  
 DR  
 XX Identifying a candidate polymorphic repeat within a coding sequence, for  
 PT understanding or treating genetic disease, comprises detecting tandem  
 PT repeats in a target coding sequence and scoring the repeats for  
 PT polymorphic probability.  
 PT  
 XX Example; Col 1163; 588pp; English.  
 PS  
 XX The invention discloses a method for identifying a candidate polymorphic  
 CC repeat within a coding sequence (expressed sequence tag, EST), which  
 CC comprises detecting tandem repeats in a target coding sequence, scoring  
 CC the repeats for polymorphic probability and generating a dataset  
 CC correlating the repeats with polymorphic probability to identify a  
 CC candidate polymorphic repeat. The computational methods (polymorphic  
 CC marker prediction of ubiquitous simple sequences, POMPOUS, and Rep-X) are  
 CC useful for identifying and detecting candidate polymorphic repeats in  
 CC human genes, which can be used to understand, treat or eliminate genetic  
 CC diseases, predispositions or adverse drug-treatment reactions. Examples  
 CC of diseases linked to nucleotide repeats are Machado-Joseph, Haw River  
 CC syndrome, Huntington's disease, fragile-X syndrome, Predreich's ataxia,  
 CC myotonic dystrophy, hyperandrogenaemia, spinal and bulbar atrophy and  
 CC spinocerebellar ataxia. The sequences presented in ABX79676-ABX80022 are  
 CC the polymorphic repeats identified for a search of human ESTs  
 XX  
 SQ Sequence 30 BP; 1 A; 9 C; 20 G; 0 T; 0 U; 0 Other;  
 Query Match 1.8%; Score 24; DB 7; Length 30;  
 Best Local Similarity 100.0%; Pred. No. 1.7;  
 Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

```
QY 56 GCGCGCGCGCGCGCGCGCGCGCGGGA 79
DB 7 GCGCGCGCGCGCGCGCGCGCGCGGGA 30

RESULT 4
ABX13592
ID ABX13592 standard; DNA; 50 BP.
AC
XX
XX
DT 10-FEB-2003 (first entry)
XX
XX
DE Human serine protein kinase KSE336 promoter fragment #8.
XX
XX Human; promoter; ds; chromosome 11p15.5-pter; astrocytoma; meningioma;
XX pancreatic adenocarcinoma; insulin-dependent diabetes mellitus 2;
XX helicoid peripapillary chorioretinal degeneration; brain; pancreas;
XX Beckwith-Wiedemann syndrome; congenital hyperinsulinism; KSE336.
XX
XX Homo sapiens.
XX
XX US6455292-B1.
XX
XX 24-SEP-2002.
XX
XX 16-AUG-2001; 2001US-00930181.
XX
XX 16-AUG-2001; 2001US-00930181.
XX
XX (ORIG-) ORIGENE TECHNOLOGIES INC.
XX
XX Shu Y, Fan W, Kovacs KF, Zidanic M, Jay G;
XX WPI; 2003-066233/06.
XX
XX New isolated polynucleotide coding without interruption for a human
XX KSE336 polypeptide useful for preventing or treating diseases/conditions
XX relating to brain and pancreas, e.g. meningioma, insulin-dependent
XX diabetes mellitus 2.
XX
XX Disclosure; Col 55; 34pp; English.
XX
XX The invention relates to an isolated polynucleotide (its complement or a
XX sequence 9% similar to it) coding without interruption for a human
XX KSE336 polypeptide, a serine protein kinase, comprising the KSE336-1 and
XX KSE336-2 splice variants appearing as ABG72382 and ABG72383. Also
XX included is a method of identifying an agent that modulates the
XX expression of KSE336 in brain, pancreas, brain progenitor or pancreas
XX progenitor cells comprising: (a) contacting a cell population comprising
XX the cells with a test agent under conditions effective for the test agent
XX to modulate the expression of KSE336; and (b) determining if the test
XX agent modulates the expression of KSE336. The polynucleotides are useful
XX as molecular targets or drug targets, and for detecting, diagnosing,
XX staging, monitoring, prognosticating, preventing or treating diseases or
XX conditions relating to brain and pancreas, such as astrocytoma,
XX meningioma, pancreatic adenocarcinoma, insulin-dependent diabetes
XX mellitus 2, helicoid peripapillary chorioretinal degeneration, Beckwith-
XX Wiedemann syndrome or congenital hyperinsulinism. The method and
XX polynucleotides are useful in research, diagnosis, drug discovery,
XX therapy, clinical medicine, forensic science and pathology. The gene for
XX KSE336 is located on chromosome 11p15.5-pter. The present sequence is
XX promoter fragment of the KSE366 gene
XX
SQ Sequence 50 BP; 2 A; 17 C; 30 G; 1 T; 0 U; 0 Other;
Query Match 1.8%; Score 24; DB 7; Length 50;
Best Local Similarity 100.0%; Pred. No. 1.7;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGCGGGA 79
DB 23 GCGCGCGCGCGCGCGCGCGCGCGGGA 46

RESULT 6
AAQ55856
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ID AAQ55856 standard; DNA; 25 BP.
XX AC AAQ55856;
XX DT 25-MAR-2003 (revised)
XX DT 25-JUL-1994 (first entry)
XX DE Fragile X probe.
XX FC; foetal cells; marker; probe; hybridise; denature; dye; label;
XX KW fluorescent; kit; detection; haemoglobin; rhesus; gamma globulin; NR;
XX KW nitrogen reductase; ss.
XX OS Homo sapiens.
XX PN WO9402646-A1.
XX PD 03-FEB-1994.
XX PF 19-JUL-1993; 93WO-US006828.
XX PR 17-JUL-1992; 92US-00915965.
XX PA (RERE-) RES DEV FOUND.
XX PI Aagari M, Prashad N, Cubbage ML, Ju S, Blick M, Bresser J;
XX WPI; 1994-048903/06.
XX Identifying foetal cells, conc. from maternal blood, using specific
XX PT marker - e.g. surface antigen, before in situ hybridisation of target
XX PT nucleic acid to detect viral infection, genetic abnormality, etc.
XX PS Disclosure; Page 73; 109pp; English.
XX CC Probes (AAQ55857-873) detect regions of 3 fragments of the HUMGLN gene
XX CC (AAQ64058). Bases 1-91 correspond to bases 2179-2269 of HUMGLN, bases 92
XX CC -314 are from 2393-2615 of HUMGLN and bases 315-443 are from 3502-3630
XX CC of HUMGLN. The probes (AAQ55854-55) were used as control, positive and
XX CC negative genetic testing probes. Probe (AAQ55856) was used to detect the
XX CC fragile X condition (Example 14) (Updated on 25-MAR-2003 to correct PN
XX CC field.)
XX SQ Sequence 25 BP; 0 A; 9 C; 16 G; 0 T; 0 U; 0 Other;

Query Match 1.7%; Score 23; DB 2; Length 25;
Best Local Similarity 100.0%; Pred. No. 4.9;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGG 78
DB 2 GCGCGCGCGCGCGCGCGCGG 24

RESULT 7
AAQ85271
ID AAQ85271 standard; DNA; 25 BP.
XX AC AAQ85271;
XX DT 25-MAR-2003 (revised)
XX DT 24-AUG-1995 (first entry)
XX DE Probe for Fragile X condition.
XX KW Prenatal diagnosis; fragile X; probe; ss.
XX OS Synthetic.
XX PN WO9503431-A1.
XX PD 02-FEB-1995.
XX

Query Match 1.7%; Score 23; DB 2; Length 25;
Best Local Similarity 100.0%; Pred. No. 4.9;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGG 78
DB 2 GCGCGCGCGCGCGCGCGCGG 24

RESULT 8
AAQ5267
ID AAQ5267 standard; DNA; 25 BP.
XX AC AAQ5267;
XX DT 14-APR-1999 (first entry)
XX DE Fragile X chromosome detecting probe.
XX KW Genetic testing; foetal cell; maternal; blood; pregnant; hybridisation;
XX KW detection; HIV, hepatitis virus; herpes virus; chromosomal abnormality;
XX KW probe; ss.
XX OS Synthetic.
XX OS Homo sapiens.
XX PN US958649-A.
XX PD 12-JAN-1999.
XX PF 31-DEC-1996; 96US-00775609.
XX PR 17-JUL-1992; 92US-00915765.
XX PR 19-JUL-1993; 93US-00094710.
XX PR 19-JUL-1994; 94WO-US008342.
XX PR 17-JAN-1995; 95US-00374144.
XX PA (APRO-) APROGENEX INC.
XX PI Blick M, Cubbage ML, Bresser J, Prashad N, Aagari M;
XX WPI; 1999-152096/13.
XX Method for distinguishing foetal cells from adult cells in blood - based
XX PT on amplification and detection of mRNA selectively expressed in foetal
XX PT cells.

```

PS Example 4, 14; Col 49; 49pp; English.

XX The invention relates to a method of enriching foetal cells from maternal  
 CC blood and for identifying such foetal cells. Foetal cells can be  
 CC distinguished from adult cells in a blood specimen by (a) treating a  
 CC blood specimen from a pregnant female to yield a mixture of cells  
 CC comprising foetal cells and adult cells; (b) amplifying one or more mRNAs  
 CC within the cells, the mRNAs being selectively expressed in target foetal  
 CC cells to be distinguished but not expressed in adult blood cells; (c)  
 CC performing in situ hybridisation on the cells under hybridising  
 CC conditions suitable to maintain cell membranes in a substantially intact  
 CC state and with a hybridisation medium comprising a detectably labelled  
 CC probe complementary to the amplified mRNA that is selectively expressed  
 CC in the target foetal cells but not expressed in adult blood cells; (d)  
 CC removing the hybridisation medium and unhybridised probe from the mixture  
 CC of cells to yield hybridised cells; and (e) detecting the labelled probe  
 CC remaining in the hybridised cells; whereby cells in which the labelled  
 CC probe is detected are identified as the target foetal cells; A second  
 CC method for determining the presence of a target nucleotide sequence in  
 CC individual foetal cells present in a cellular specimen is also provided.  
 CC The methods (especially the second) is useful for detecting HIV,  
 CC hepatitis viruses or herpes viruses in foetal cells, or for detecting  
 CC chromosomal abnormalities in foetal cells. The present sequence  
 CC represents a probe used for the detection of the Fragile X chromosome in  
 CC amniocytes and in peripheral blood mononuclear cells

XX SQ Sequence 25 BP; 0 A; 9 C; 16 G; 0 T; 0 U; 0 Other;

Query Match 1.7%; Score 23; DB 2; Length 25;  
 Best Local Similarity 100.0%; Pred. No. 4.9;  
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
 DB 2 GCGCGCGCGCGCGCGCGCGCGG 24

RESULT 9

ID AAC60682/C  
 XX AAC60682 standard; DNA; 45 BP.

AC AAC60682;

XX 29-JAN-2001 (first entry)

XX Triplet repeat block exemplary test sequence #3.

XX Mismatch-binding protein; MutS; hexA; MBP; hMSH2; forensic medicine;  
 KW disease diagnosis; cancer; epidemiology; allele identification;  
 KW infectious agent; triplet repeat; PCR primer; probe; ss.

XX Unidentified.

XX US6120992-A.

XX 19-SEP-2000.

XX 04-MAR-1996; 96US-00608016.

XX 04-NOV-1993; 93US-00147785.

XX 28-APR-1995; 95US-00431081.

XX (VALI-) VALIGENE CORP.

XX Wagner RE;

XX WPI; 2000-601481/57.

XX Use of immobilized MutS protein that binds to nucleic acid hybrid with  
 PT single base mismatch, for detecting in test DNA sample from diseased  
 PT human, presence of heteroduplex having deletion or addition 4  
 XX nucleotides.

PS Disclosure; Fig 12a; 62pp; English.

XX The present invention is concerned with the detection of DNA sequences  
 CC which may have a single mismatched base or up to 4 bases inserted or  
 CC deleted from the sequence when compared with the wild-type. The E. coli  
 CC MutS protein or its homologues (for example hexA and hMSH2) can be used  
 CC to bind to a sample containing the nucleic acid sequence of interest and  
 CC a wild-type sequence. If there is a mismatch, the MutS protein will bind  
 CC to the duplex and can then be detected. This method can be used in the  
 CC diagnosis of diseases such as cancer and those caused by triplet repeat  
 CC expansions, including Fragile X syndrome, myotonic dystrophy,  
 CC Huntington's disease, spino-cerebellar ataxia type 1, spinal bulbar  
 CC muscular atrophy, Machado-Joseph disease and dentatorubralpallidoluysian  
 CC atrophy. It can also be used to detect and determine the epidemiology of  
 CC infectious diseases, in specific allele identification and to detect  
 CC mutations and polymorphisms. The sequences shown in AAC60681-C60683 were  
 CC all used to demonstrate the method

XX SQ Sequence 45 BP; 0 A; 18 C; 9 G; 0 T; 0 U; 18 Other;

Query Match 1.7%; Score 23; DB 3; Length 45;  
 Best Local Similarity 100.0%; Pred. No. 4.7;  
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
 DB 35 GCGCGCGCGCGCGCGCGCGCGG 13

RESULT 10

AAD40340/C

ID AAD40340 standard; DNA; 45 BP.

XX AAD40340;

XX 22-OCT-2002 (first entry)

XX Test DNA #3 used in the triplet repeat diagnostic method.

XX Mismatch binding protein; heteroduplex DNA; triplet repeat block; TRB;  
 KW fragile X syndrome; myotonic dystrophy; Huntington's disease;  
 KW spino-cerebellar ataxia type I; spinal bulbar muscular atrophy;  
 KW Machado-Joseph disease; dentatorubralpallidoluysian atrophy; ss.

XX Unidentified.

XX US6323147-B1.

XX 11-DEC-2001.

XX 04-FEB-2000; 2000US-00497933.

XX 04-NOV-1993; 93US-00147785.

XX 28-APR-1995; 95US-00431081.

XX 04-MAR-1996; 96US-00608016.

XX (VALI-) VALIGEN US INC.

XX Wagner RE;

XX WPI; 2002-187350/24.

XX Determining mismatch binding protein in sample comprises mixing sample  
 PT with labeled heteroduplex DNA, contacting mixture with immobilized  
 PT mismatch binding protein and detecting amount of DNA bound and comparing  
 PT with control.

XX Disclosure; Fig 12B; 62pp; English.

XX The invention relates to a method for determining the presence of a  
 CC functional mismatch binding protein. The method comprising mixing  
 CC biological fluid sample with detectably labelled heteroduplex DNA to form  
 CC a mixture, contacting the mixture with immobilised mismatch-binding

CC protein, detecting the binding of the DNA to the protein and comparing  
 CC the amount of the heteroduplex DNA bound with the amount of the DNA bound  
 CC when control sample is utilised instead of fluid sample. The method is  
 CC useful for determining the presence of a functional mismatch-binding  
 CC protein in a biological fluid sample preferably presence of a repeat  
 CC block especially a triplet repeat block (TRB) of unit sequence 5'-(CAG)<sub>n</sub>  
 CC or 5'-(CTG)<sub>n</sub> 3'-(GAC)<sub>n</sub> in test DNA in a sample preferably by  
 CC a competitive assay method, where TRB is longer (or shorter) than TRB in  
 CC a diagnostic oligonucleotide probe, and n is an integer and is number of  
 CC repeats of TRB in the test DNA. The method is useful for diagnosing a  
 CC variety of disease states or susceptibilities such as fragile X syndrome,  
 CC myotonic dystrophy, Huntington's disease, spinocerebellar ataxia type 1,  
 CC spinal bulbar muscular atrophy, Machado-Joseph disease and  
 CC dentatorubralpallidolysian atrophy. The present sequence is test DNA  
 CC used in the triplet repeat diagnostic method  
 XX  
 SQ Sequence 45 BP; 0 A; 18 C; 9 G; 0 T; 0 U; 18 Other;

Query Match 1.7%; Score 23; DB 6; Length 45;  
 Best Local Similarity 100.0%; Pred. No. 4.7;  
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
 DB 35 GCGCGCGCGCGCGCGCGCGCGG 13

## RESULT 11

ABS67036  
 ID ABS67036 standard; DNA; 47 BP.

AC ABS67036;

DT 29-NOV-2002 (first entry)

DE Human MRP-1 polymorphic DNA region #298.

XX Human; multidrug resistance-associated protein 1; MRP-1; ss; cancer;  
 XX renal cancer; cytostatic; single nucleotide polymorphism.

OS Homo sapiens;

XX WO200259142-A2.

PD 01-AUG-2002.

PF 25-JAN-2002; 2002WO-EP000796.

PR 26-JAN-2001; 2001EP-00101651.

PA (EPID-) EPIDAUROS BIOTECHNOLOGIES AG.

PI Brinkmann U, Hoffmeyer S, Mornhinweg E;

XX WPI; 2002-657475/70.

XX Novel multidrug resistance-associated protein 1 polynucleotide useful for  
 PT diagnosis and treatment of cancer and multidrug resistance related  
 PT diseases, and for identifying single nucleotide polymorphisms.

PS Claim 1; Page 85; 198pp; English.

XX The invention relates to a multidrug resistance-associated protein 1 (MRP  
 CC -1) polynucleotide. The polynucleotide is useful in an in vitro method  
 CC for identifying a single nucleotide polymorphism and for identifying and  
 CC obtaining a pro-drug or drug capable of modulating the activity of a  
 CC molecular variant of MRP-1 or for identifying and obtaining an inhibitor  
 CC of the activity of a molecular variant of MRP-1. The sequences are useful  
 CC for diagnosing a disorder related to the presence of a molecular variant  
 CC of MRP-1 or susceptibility to such a disorder, where the disorder is  
 CC cancer (particularly renal cancer) or a disease related to multidrug  
 CC resistance. This sequence represents a human MRP-1 polymorphic DNA region

XX

SQ Sequence 47 BP; 3 A; 15 C; 27 G; 2 T; 0 U; 0 Other;

Query Match 1.7%; Score 23; DB 6; Length 47;  
 Best Local Similarity 100.0%; Pred. No. 4.7;  
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
 DB 11 GCGCGCGCGCGCGCGCGCGCGG 33

## RESULT 12

ABS67035/C

ID ABS67035 standard; DNA; 47 BP.

XX ABS67035;

AC ABS67035;

DT 29-NOV-2002 (first entry)

DE Human MRP-1 polymorphic DNA region #297.

XX Human; multidrug resistance-associated protein 1; MRP-1; ss; cancer;  
 XX renal cancer; cytostatic; single nucleotide polymorphism.

OS Homo sapiens.

XX WO200259142-A2.

PD 01-AUG-2002.

PF 25-JAN-2002; 2002WO-EP000796.

PR 26-JAN-2001; 2001EP-00101651.

PA (EPID-) EPIDAUROS BIOTECHNOLOGIES AG.

PI Brinkmann U, Hoffmeyer S, Mornhinweg E;

XX WPI; 2002-657475/70.

XX Novel multidrug resistance-associated protein 1 polynucleotide useful for  
 PT diagnosis and treatment of cancer and multidrug resistance related  
 PT diseases, and for identifying single nucleotide polymorphisms.

PS Claim 1; Page 85; 198pp; English.

XX The invention relates to a multidrug resistance-associated protein 1 (MRP  
 CC -1) polynucleotide. The polynucleotide is useful in an in vitro method  
 CC for identifying a single nucleotide polymorphism and for identifying and  
 CC obtaining a pro-drug or drug capable of modulating the activity of a  
 CC molecular variant of MRP-1 or for identifying and obtaining an inhibitor  
 CC of the activity of a molecular variant of MRP-1. The sequences are useful  
 CC for diagnosing a disorder related to the presence of a molecular variant  
 CC of MRP-1 or susceptibility to such a disorder, where the disorder is  
 CC cancer (particularly renal cancer) or a disease related to multidrug  
 CC resistance. This sequence represents a human MRP-1 polymorphic DNA region

SQ Sequence 47 BP; 2 A; 27 C; 15 G; 3 T; 0 U; 0 Other;

Query Match 1.7%; Score 23; DB 6; Length 47;  
 Best Local Similarity 100.0%; Pred. No. 4.7;  
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
 DB 37 GCGCGCGCGCGCGCGCGCGCGG 15

## RESULT 13

AAC60679

ID AAC60679 standard; DNA; 48 BP.

XX AAC60679;

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XX 29-JAN-2001 (first entry)
XX Triplet repeat block exemplary test sequence #1.
XX Mismatch-binding protein; Muts; hexA; MBP; hMSH2; forensic medicine;
XX disease diagnosis; cancer; epidemiology; allele identification;
XX infectious agent; triplet repeat; PCR primer; probe; ss.
XX Unidentified.
XX OS
XX US6120992-A.
XX PN
XX 19-SEP-2000.
XX PD
XX 04-MAR-1996; 96US-00608016.
XX PF
XX 04-NOV-1993; 93US-00147785.
XX PR
XX 28-APR-1995; 95US-00431081.
XX ER
XX (VALI-) VALIGENE CORP.
XX PA
XX Wagner RE;
XX PI
XX WPI; 2000-601481/57.
XX DR
XX
XX Use of immobilized Muts protein that binds to nucleic acid hybrid with
XX single base mismatch, for detecting in test DNA sample from diseased
XX human, presence of heteroduplex having deletion or addition 4
XX nucleotides.
XX PT
XX Disclosure; Fig 12a; 62pp; English.
XX PS
XX The present invention is concerned with the detection of DNA sequences
XX which may have a single mismatched base or up to 4 bases inserted or
XX deleted from the sequence when compared with the wild-type. The E. coli
XX Muts protein or its homologues (for example hexA and hMSH2) can be used
XX to bind to a sample containing the nucleic acid sequence of interest and
XX a wild-type sequence. If there is a mismatch, the Muts protein will bind
XX to the duplex and can then be detected. This method can be used in the
XX diagnosis of diseases such as cancer and those caused by triplet repeat
XX expansions, including Fragile X syndrome, myotonic dystrophy.
XX CC
XX Huntington's disease, spino-cerebellar ataxia type 1, spinal bulbar
XX muscular atrophy, Machado-Joseph disease and dentatorubralpallidoluysian
XX atrophy. It can also be used to detect and determine the epidemiology of
XX infectious diseases, in specific allele identification and to detect
XX mutations and polymorphisms. The sequences shown in AAC60661-C60683 were
XX all used to demonstrate the method
XX SQ
XX Sequence 48 BP; 0 A; 10 C; 20 G; 0 T; 0 U; 18 Other;
XX
XX Query Match 1.7%; Score 23; DB 3; Length 48;
XX Best Local Similarity 100.0%; Pred. No. 4.7;
XX Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
XX QY 56 GCGCGCGCGCGCGCGCGCGCGCGG 78
XX |||||
XX DB 10 GCGCGCGCGCGCGCGCGCGCGG 32
XX |||||
XX
XX RESULT 14
XX AAD40337
XX ID AAD40337 standard; DNA; 48 BP.
XX XX
XX AC AAD40337;
XX XX
XX 22-OCT-2002 (first entry)
XX DT
XX Test DNA #1 used in the triplet repeat diagnostic method.
XX DE
XX Mismatch binding protein; heteroduplex DNA; triplet repeat block; TRB;
XX fragile X syndrome; myotonic dystrophy; Huntington's disease;
XX spino-cerebellar ataxia type I; spinal bulbar muscular atrophy;
XX KW

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KW Machado-Joseph disease; dentatorubralpallidoluysian atrophy; ss.
XX Unidentified.
XX OS
XX US6329147-B1.
XX PN
XX 11-DEC-2001.
XX PD
XX 04-FEB-2000; 2000US-00497933.
XX PF
XX 04-NOV-1993; 93US-00147785.
XX PR
XX 28-APR-1995; 95US-00431081.
XX ER
XX 04-MAR-1996; 96US-00608016.
XX PA
XX (VALI-) VALIGENE US INC.
XX PI
XX Wagner RE;
XX DR
XX WPI; 2002-187350/24.
XX XX
XX Determining mismatch binding protein in sample comprises mixing sample
XX with labeled heteroduplex DNA, contacting mixture with immobilized
XX mismatch binding protein and detecting amount of DNA bound and comparing
XX with control.
XX PT
XX Disclosure; Fig 12a; 62pp; English.
XX PS
XX The invention relates to a method for determining the presence of a
XX functional mismatch binding protein. The method comprising mixing
XX biological fluid sample with detectably labelled heteroduplex DNA to form
XX a mixture, contacting the mixture with immobilised mismatch-binding
XX protein, detecting the binding of the DNA to the protein and comparing
XX the amount of the heteroduplex DNA bound with the amount of the DNA bound
XX when control sample is utilised instead of fluid sample. The method is
XX useful for determining the presence of a functional mismatch-binding
XX protein in a biological fluid sample preferably presence of a repeat
XX block especially a triplet repeat block (TRB) of unit sequence 5'-(CGG)n
XX or 5'-(CTG)n 3'-(GCC)n 3'-(GAC)n in test DNA in a sample preferably by
XX a competitive assay method, where TRB is longer (or shorter) than TRB in
XX a diagnostic oligonucleotide probe, and n is an integer and is number of
XX repeats of TRB in the test DNA. The method is useful for diagnosing a
XX variety of disease states or susceptibilities such as fragile X syndrome,
XX myotonic dystrophy, Huntington's disease, spino-cerebellar ataxia type I,
XX spinal bulbar muscular atrophy, Machado-Joseph disease and
XX dentatorubralpallidoluysian atrophy. The present sequence is test DNA
XX used in the triplet repeat diagnostic method
XX CC
XX SQ
XX Sequence 48 BP; 0 A; 10 C; 20 G; 0 T; 0 U; 18 Other;
XX
XX Query Match 1.7%; Score 23; DB 6; Length 48;
XX Best Local Similarity 100.0%; Pred. No. 4.7;
XX Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
XX QY 56 GCGCGCGCGCGCGCGCGCGCGCGG 78
XX |||||
XX DB 11 GCGCGCGCGCGCGCGCGCGCGG 33
XX |||||
XX
XX RESULT 15
XX ABS67037/c
XX ID ABS67037 standard; DNA; 48 BP.
XX XX
XX AC ABS67037;
XX XX
XX 29-NOV-2002 (first entry)
XX DT
XX Human MRP-1 polymorphic DNA region #299.
XX DE
XX Human; multidrug resistance-associated protein 1; MRP-1; ss; cancer;
XX renal cancer; cytostatic; single nucleotide polymorphism.
XX KW
XX Homo sapiens.
XX XX

```

PN WO200259142-A2.  
 XX  
 PD 01-AUG-2002.  
 XX  
 PF 25-JAN-2002; 2002WO-EP000796.  
 XX  
 PR 26-JAN-2001; 2001EP-00101651.  
 XX  
 PA (EPID-) EPIDAUROS BIOTECHNOLOGIES AG.  
 XX  
 PI Brinkmann U, Hoffmeyer S, Mornhinweg E;  
 XX  
 DR WPI; 2002-657475/70.  
 XX  
 PT Novel multidrug resistance-associated protein 1 polynucleotide useful for  
 PT diagnosis and treatment of cancer and multidrug resistance related  
 PT diseases, and for identifying single nucleotide polymorphisms.  
 XX  
 PS Example 9; Page 85; 198pp; English.  
 XX  
 CC The invention relates to a multidrug resistance-associated protein 1 (MRP  
 CC -1) polynucleotide. The polynucleotide is useful in an in vitro method  
 CC for identifying a single nucleotide polymorphism and for identifying and  
 CC obtaining a pro-drug or drug capable of modulating the activity of a  
 CC molecular variant of MRP-1 or for identifying and obtaining an inhibitor  
 CC of the activity of a molecular variant of MRP-1. The sequences are useful  
 CC for diagnosing a disorder related to the presence of a molecular variant  
 CC of MRP-1 or susceptibility to such a disorder, where the disorder is  
 CC cancer (particularly renal cancer) or a disease related to multidrug  
 CC resistance. This sequence represents a human MRP-1 polymorphic DNA region  
 XX  
 SQ Sequence 48 BP; 2 A; 27 C; 15 G; 3 T; 0 U; 1 Other;  
 Query Match 1.7%; Score 23; DB 6; Length 48;  
 Best Local Similarity 100.0%; Pred. No. 4.7;  
 Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 Qy 56 GGCGGCGGCGGCGGCGGCGGCGG 78  
 Db 37 GGCGGCGGCGGCGGCGGCGGCGG 15

Search completed: July 1, 2004, 19:19:07  
 Job time : 582 secs

GenCore version 5.1.6  
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OM nucleic - nucleic search, using sw model  
Run on: July 1, 2004, 20:48:10 ; Search time 644 Seconds  
(without alignments)  
9895.618 Million cell updates/sec

Title: US-09-934-249-1  
Perfect score: 1321  
Sequence: 1 cgacgcggtctcgagcga.....ctgcgttagtgaaaggcag 1321

Scoring table: OLIGO\_NUC  
Gapop 60.0 , Gapext 60.0

Searched: 3163042 seqs, 2412103800 residues

Word size : 0

Total number of hits satisfying chosen parameters: 1726952

Minimum DB seq length: 0

Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database : Published Applications NA:\*

- 1: /cgn2\_6/ptodata/1/pubna/US07\_PUBCOMB.seq:\*
- 2: /cgn2\_6/ptodata/1/pubna/PCT\_NEW\_PUB.seq:\*
- 3: /cgn2\_6/ptodata/1/pubna/US06\_PUBCOMB.seq:\*
- 4: /cgn2\_6/ptodata/1/pubna/US06\_PUBCOMB.seq:\*
- 5: /cgn2\_6/ptodata/1/pubna/US07\_NEW\_PUB.seq:\*
- 6: /cgn2\_6/ptodata/1/pubna/PCTUS\_PUBCOMB.seq:\*
- 7: /cgn2\_6/ptodata/1/pubna/US08\_NEW\_PUB.seq:\*
- 8: /cgn2\_6/ptodata/1/pubna/US08\_PUBCOMB.seq:\*
- 9: /cgn2\_6/ptodata/1/pubna/US09A\_PUBCOMB.seq:\*
- 10: /cgn2\_6/ptodata/1/pubna/US09B\_PUBCOMB.seq:\*
- 11: /cgn2\_6/ptodata/1/pubna/US09C\_PUBCOMB.seq:\*
- 12: /cgn2\_6/ptodata/1/pubna/US09\_NEW\_PUB.seq:\*
- 13: /cgn2\_6/ptodata/1/pubna/US09\_NEW\_PUB.seq:\*
- 14: /cgn2\_6/ptodata/1/pubna/US10A\_PUBCOMB.seq:\*
- 15: /cgn2\_6/ptodata/1/pubna/US10B\_PUBCOMB.seq:\*
- 16: /cgn2\_6/ptodata/1/pubna/US10C\_PUBCOMB.seq:\*
- 17: /cgn2\_6/ptodata/1/pubna/US10\_NEW\_PUB.seq:\*
- 18: /cgn2\_6/ptodata/1/pubna/US60\_NEW\_PUB.seq:\*
- 19: /cgn2\_6/ptodata/1/pubna/US60\_PUBCOMB.seq:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

# SUMMARIES

| Result No. | Score | Query Match | Length | ID | Description       |
|------------|-------|-------------|--------|----|-------------------|
| 1          | 24    | 1.8         | 50     | 15 | US-10-195-072-14  |
| 2          | 24    | 1.8         | 50     | 15 | US-10-195-071-14  |
| 3          | 23    | 1.7         | 50     | 15 | US-10-434-479-78  |
| 4          | 23    | 1.7         | 50     | 15 | US-10-126-448-2   |
| 5          | 22    | 1.7         | 22     | 15 | US-10-390-045-8   |
| 6          | 22    | 1.7         | 22     | 17 | US-10-434-479-8   |
| 7          | 22    | 1.7         | 22     | 17 | US-10-434-479-70  |
| 8          | 21    | 1.6         | 29     | 15 | US-09-828-034-10  |
| 9          | 21    | 1.6         | 39     | 15 | US-10-157-382-7   |
| 10         | 21    | 1.6         | 39     | 15 | US-09-888-326-410 |
| 11         | 20    | 1.5         | 20     | 10 | US-09-776-479-243 |
| 12         | 20    | 1.5         | 20     | 10 | US-09-776-479-243 |
| 13         | 20    | 1.5         | 20     | 13 | US-10-314-578-243 |
| 14         | 20    | 1.5         | 20     | 13 | US-09-776-479-243 |

|    |    |     |    |    |                   |                   |
|----|----|-----|----|----|-------------------|-------------------|
| 15 | 20 | 1.5 | 20 | 15 | US-10-112-653-235 | Sequence 235, App |
| 16 | 20 | 1.5 | 20 | 15 | US-10-017-995-243 | Sequence 243, App |
| 17 | 20 | 1.5 | 20 | 15 | US-10-390-045-9   | Sequence 9, Appli |
| 18 | 20 | 1.5 | 20 | 17 | US-10-434-479-9   | Sequence 9, Appli |
| 19 | 20 | 1.5 | 50 | 15 | US-10-195-072-13  | Sequence 13, Appl |
| 20 | 20 | 1.5 | 50 | 15 | US-10-195-071-13  | Sequence 13, Appl |
| 21 | 19 | 1.4 | 25 | 17 | US-10-434-479-72  | Sequence 72, Appl |
| 22 | 19 | 1.4 | 26 | 13 | US-10-236-392-385 | Sequence 385, App |
| 23 | 19 | 1.4 | 28 | 15 | US-10-170-663-6   | Sequence 6, Appli |
| 24 | 19 | 1.4 | 48 | 15 | US-10-205-942-9   | Sequence 9, Appli |
| 25 | 19 | 1.4 | 48 | 15 | US-10-205-942-10  | Sequence 10, Appl |
| 26 | 18 | 1.4 | 18 | 10 | US-09-500-700-68  | Sequence 68, Appl |
| 27 | 18 | 1.4 | 18 | 10 | US-10-314-405-45  | Sequence 45, Appl |
| 28 | 18 | 1.4 | 18 | 17 | US-10-627-473-45  | Sequence 45, Appl |
| 29 | 18 | 1.4 | 20 | 10 | US-09-888-326-410 | Sequence 410, App |
| 30 | 18 | 1.4 | 20 | 10 | US-09-776-479-243 | Sequence 243, App |
| 31 | 18 | 1.4 | 20 | 13 | US-10-314-578-243 | Sequence 243, App |
| 32 | 18 | 1.4 | 20 | 13 | US-09-776-479-243 | Sequence 243, App |
| 33 | 18 | 1.4 | 20 | 15 | US-10-112-653-235 | Sequence 235, App |
| 34 | 18 | 1.4 | 20 | 15 | US-10-017-995-243 | Sequence 243, App |
| 35 | 18 | 1.4 | 21 | 9  | US-09-828-034-10  | Sequence 10, Appl |
| 36 | 18 | 1.4 | 27 | 9  | US-09-848-213-18  | Sequence 18, Appl |
| 37 | 18 | 1.4 | 27 | 9  | US-09-848-213-19  | Sequence 19, Appl |
| 38 | 18 | 1.4 | 28 | 10 | US-09-500-700-38  | Sequence 38, Appl |
| 39 | 18 | 1.4 | 28 | 15 | US-10-422-934-21  | Sequence 21, Appl |
| 40 | 18 | 1.4 | 30 | 9  | US-09-973-145-11  | Sequence 11, Appl |
| 41 | 18 | 1.4 | 30 | 15 | US-10-174-368A-11 | Sequence 11, Appl |
| 42 | 18 | 1.4 | 30 | 15 | US-10-264-137-12  | Sequence 12, Appl |
| 43 | 18 | 1.4 | 30 | 16 | US-10-339-712-12  | Sequence 12, Appl |
| 44 | 18 | 1.4 | 30 | 17 | US-10-642-763-19  | Sequence 19, Appl |
| 45 | 18 | 1.4 | 30 | 17 | US-10-642-763-21  | Sequence 21, Appl |

## ALIGNMENTS

### RESULT 1

US-10-195-072-14  
; Sequence 14, Application US/10195072  
; Publication No. US20030092036A1  
; GENERAL INFORMATION:  
; APPLICANT: Origene Technologies  
; TITLE OF INVENTION: Full-length Serine Protein Kinase in Brain and Pancreas  
; FILE REFERENCE: 16U 101 C2  
; CURRENT APPLICATION NUMBER: US/10/195,072  
; CURRENT FILING DATE: 2002-07-15  
; PRIOR APPLICATION NUMBER: US 09/930,181  
; PRIOR FILING DATE: 2001-08-16  
; NUMBER OF SEQ ID NOS: 18  
; SOFTWARE: PatentIn version 3.1  
; SEQ ID NO 14  
; LENGTH: 50  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-10-195-072-14

Query Match 1.8%; Score 24; DB 15; Length 50;  
Best Local Similarity 100.0%; Pred. No. 0.41;  
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Cy 56 GCGCGCGCGCGCGCGCGCGCGCGA 79  
Db 23 GCGCGCGCGCGCGCGCGCGCGA 46

### RESULT 2

US-10-195-071-14  
; Sequence 14, Application US/10195071  
; Publication No. US2003009271A1  
; GENERAL INFORMATION:  
; APPLICANT: Origene Technologies  
; TITLE OF INVENTION: Full-length Serine Protein Kinase in Brain and Pancreas  
; FILE REFERENCE: 16U 101 C1

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; CURRENT APPLICATION NUMBER: US/10/195,071
; CURRENT FILING DATE: 2002-07-15
; PRIOR APPLICATION NUMBER: US 09/930,181
; PRIOR FILING DATE: 2001-08-16
; NUMBER OF SEQ ID NOS: 18
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 14
; LENGTH: 50
; TYPE: DNA
; ORGANISM: Homo sapiens
US-10-195-071-14

Query Match      1.8%; Score 24; DB 15; Length 50;
Best Local Similarity 100.0%; Pred. No. 0.41;
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGGA 79
Db 23 GCGCGCGCGCGCGCGCGCGCGGA 46

RESULT 3
US-10-434-479-78/c
; Sequence 78, Application US/10434479
; Publication No. US20040092469A1
; GENERAL INFORMATION:
; APPLICANT: SRIVASTAVA, SHIV
; APPLICANT: MOUL, JUDD W.
; APPLICANT: XU, LINDA L.
; TITLE OF INVENTION: ANDROGEN-REGULATED PMEPA1 GENE AND POLYPEPTIDES
; FILE REFERENCE: 04995.0057-02000
; CURRENT APPLICATION NUMBER: US/10/434,479
; CURRENT FILING DATE: 2003-05-09
; PRIOR APPLICATION NUMBER: 10/390,045
; PRIOR FILING DATE: 2003-03-18
; PRIOR APPLICATION NUMBER: 03/769,482
; PRIOR FILING DATE: 2001-01-26
; PRIOR APPLICATION NUMBER: 60/178,772
; PRIOR FILING DATE: 2000-01-28
; PRIOR APPLICATION NUMBER: 60/179,045
; PRIOR FILING DATE: 2000-01-31
; NUMBER OF SEQ ID NOS: 81
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 78
; LENGTH: 23
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Primer
US-10-434-479-78

Query Match      1.7%; Score 23; DB 17; Length 23;
Best Local Similarity 100.0%; Pred. No. 1.4;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 523 GGAGCTGGAGTTTTCAGATCA 545
Db 23 GGAGCTGGAGTTTTCAGATCA 1

RESULT 4
US-10-126-448-2/c
; Sequence 2, Application US/10126448
; Publication No. US20030148422A1
; GENERAL INFORMATION:
; APPLICANT: Doring, Volker
; APPLICANT: Nangle, Leslie A.
; APPLICANT: Hendrickson, Tamara L.
; APPLICANT: de Crecy-Lagard, Valerie
; APPLICANT: Schimmel, Paul
; APPLICANT: Marliere, Philippe
; TITLE OF INVENTION: Method for Diversifying the Chemical
; Composition of Proteins Produced in vivo by Genetically
```

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; TITLE OF INVENTION: Disabling the Editing Function of their Aminoacyl tRNA
; TITLE OF INVENTION: Synthetases
; FILE REFERENCE: CAB1-004
; CURRENT APPLICATION NUMBER: US/10/126,448
; CURRENT FILING DATE: 2002-04-19
; PRIOR APPLICATION NUMBER: US 60/285,495
; PRIOR FILING DATE: 2001-04-19
; NUMBER OF SEQ ID NOS: 6
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 2
; LENGTH: 50
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: synthetic oligonucleotide
; FEATURE:
; NAME/KEY: misc_feature
; LOCATION: 1
; OTHER INFORMATION: N = phosphorylated cytidine
US-10-126-448-2
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Query Match      1.7%; Score 23; DB 15; Length 50;
Best Local Similarity 100.0%; Pred. No. 1.2;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
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QY 56 GCGCGCGCGCGCGCGCGCGCG 78
Db 43 GCGCGCGCGCGCGCGCGCGCG 21
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```
RESULT 5
US-10-390-045-8/c
; Sequence 8, Application US/10390045
; Publication No. US20030170713A1
; GENERAL INFORMATION:
; APPLICANT: SRIVASTAVA, SHIV
; APPLICANT: MOUL, JUDD W.
; APPLICANT: XU, LINDA L.
; APPLICANT: SEGAWA, TAKEHIKO
; TITLE OF INVENTION: PROSTATE-SPECIFIC ANDROGEN-SIGNALING-ASSOCIATED
; TITLE OF INVENTION: POYNUCLEOTIDE ARRAY
; FILE REFERENCE: 04995.0057-00000
; CURRENT APPLICATION NUMBER: US/10/390,045
; CURRENT FILING DATE: 2003-03-18
; PRIOR APPLICATION NUMBER: US/09/769,482
; PRIOR FILING DATE: 2001-01-26
; PRIOR APPLICATION NUMBER: 60/178,772
; PRIOR FILING DATE: 2000-01-28
; PRIOR APPLICATION NUMBER: 60/179,045
; PRIOR FILING DATE: 2000-01-31
; NUMBER OF SEQ ID NOS: 67
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 8
; LENGTH: 22
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Primer
US-10-390-045-8
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Query Match      1.7%; Score 22; DB 15; Length 22;
Best Local Similarity 100.0%; Pred. No. 4.3;
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
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```
QY 838 CGAGATCGACCTGCCACCCACC 859
Db 22 CGAGATCGACCTGCCACCCACC 1
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```
RESULT 6
US-10-434-479-8/c
; Sequence 8, Application US/10434479
; Publication No. US20040092469A1
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; GENERAL INFORMATION:
; APPLICANT: SRIVASTAVA, SHIV
; APPLICANT: MOUL, JUDD W.
; APPLICANT: XU, LINDA L.
; TITLE OF INVENTION: ANDROGEN-REGULATED PMPAL GENE AND POLYPEPTIDES
; FILE REFERENCE: 04995.0057-02000
; CURRENT APPLICATION NUMBER: US/10/434,479
; CURRENT FILING DATE: 2003-05-09
; PRIOR APPLICATION NUMBER: 10/390,045
; PRIOR FILING DATE: 2003-03-18
; PRIOR APPLICATION NUMBER: 09/769,482
; PRIOR FILING DATE: 2001-01-26
; PRIOR APPLICATION NUMBER: 60/178,772
; PRIOR FILING DATE: 2000-01-28
; PRIOR APPLICATION NUMBER: 60/179,045
; PRIOR FILING DATE: 2000-01-31
; NUMBER OF SEQ ID NOS: 81
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 8
; LENGTH: 22
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Primer
US-10-434-479-8

Query Match      1.7%; Score 22; DB 17; Length 22;
Best Local Similarity 100.0%; Pred. No. 4.3;
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 838 CGAGTCGACCTGCCACCCACC 859
DB 22 CGAGTCGACCTGCCACCCACC 1

RESULT 7
US-10-434-479-70/c
; Sequence 70, Application US/10434479
; Publication No. US20040092469A1
; GENERAL INFORMATION:
; APPLICANT: SRIVASTAVA, SHIV
; APPLICANT: MOUL, JUDD W.
; APPLICANT: XU, LINDA L.
; TITLE OF INVENTION: ANDROGEN-REGULATED PMPAL GENE AND POLYPEPTIDES
; FILE REFERENCE: 04995.0057-02000
; CURRENT APPLICATION NUMBER: US/10/434,479
; CURRENT FILING DATE: 2003-05-09
; PRIOR APPLICATION NUMBER: 10/390,045
; PRIOR FILING DATE: 2003-03-18
; PRIOR APPLICATION NUMBER: 09/769,482
; PRIOR FILING DATE: 2001-01-26
; PRIOR APPLICATION NUMBER: 60/178,772
; PRIOR FILING DATE: 2000-01-28
; PRIOR APPLICATION NUMBER: 60/179,045
; PRIOR FILING DATE: 2000-01-31
; NUMBER OF SEQ ID NOS: 81
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 70
; LENGTH: 22
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Primer
US-10-434-479-70

Query Match      1.7%; Score 22; DB 17; Length 22;
Best Local Similarity 100.0%; Pred. No. 4.3;
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1246 GAAGGATAAACAGAAAGGACAC 1267
DB 22 GAAGGATAAACAGAAAGGACAC 1

; GENERAL INFORMATION:
; APPLICANT: SRIVASTAVA, SHIV
; APPLICANT: MOUL, JUDD W.
; APPLICANT: XU, LINDA L.
; TITLE OF INVENTION: ANDROGEN-REGULATED PMPAL GENE AND POLYPEPTIDES
; FILE REFERENCE: 04995.0057-02000
; CURRENT APPLICATION NUMBER: US/10/434,479
; CURRENT FILING DATE: 2003-05-09
; PRIOR APPLICATION NUMBER: 10/390,045
; PRIOR FILING DATE: 2003-03-18
; PRIOR APPLICATION NUMBER: 09/769,482
; PRIOR FILING DATE: 2001-01-26
; PRIOR APPLICATION NUMBER: 60/178,772
; PRIOR FILING DATE: 2000-01-28
; PRIOR APPLICATION NUMBER: 60/179,045
; PRIOR FILING DATE: 2000-01-31
; NUMBER OF SEQ ID NOS: 81
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 8
; LENGTH: 22
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Primer
US-10-434-479-8

Query Match      1.7%; Score 22; DB 17; Length 22;
Best Local Similarity 100.0%; Pred. No. 4.3;
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 838 CGAGTCGACCTGCCACCCACC 859
DB 22 CGAGTCGACCTGCCACCCACC 1

RESULT 7
US-10-434-479-70/c
; Sequence 70, Application US/10434479
; Publication No. US20040092469A1
; GENERAL INFORMATION:
; APPLICANT: SRIVASTAVA, SHIV
; APPLICANT: MOUL, JUDD W.
; APPLICANT: XU, LINDA L.
; TITLE OF INVENTION: ANDROGEN-REGULATED PMPAL GENE AND POLYPEPTIDES
; FILE REFERENCE: 04995.0057-02000
; CURRENT APPLICATION NUMBER: US/10/434,479
; CURRENT FILING DATE: 2003-05-09
; PRIOR APPLICATION NUMBER: 10/390,045
; PRIOR FILING DATE: 2003-03-18
; PRIOR APPLICATION NUMBER: 09/769,482
; PRIOR FILING DATE: 2001-01-26
; PRIOR APPLICATION NUMBER: 60/178,772
; PRIOR FILING DATE: 2000-01-28
; PRIOR APPLICATION NUMBER: 60/179,045
; PRIOR FILING DATE: 2000-01-31
; NUMBER OF SEQ ID NOS: 81
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 70
; LENGTH: 22
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Primer
US-10-434-479-70

Query Match      1.7%; Score 22; DB 17; Length 22;
Best Local Similarity 100.0%; Pred. No. 4.3;
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1246 GAAGGATAAACAGAAAGGACAC 1267
DB 22 GAAGGATAAACAGAAAGGACAC 1
```

```

RESULT 8
US-09-828-034-10/c
; Sequence 10, Application US/09828034
; Patent No. US20020064771A1
; GENERAL INFORMATION:
; APPLICANT: Zhong, Weidong
; APPLICANT: Hong, Zhi
; APPLICANT: Ferrari, Eric
; TITLE OF INVENTION: HCV REPLICASE COMPLEXES
; FILE REFERENCE: INC1165
; CURRENT APPLICATION NUMBER: US/09/828,034
; CURRENT FILING DATE: 2001-04-06
; PRIOR APPLICATION NUMBER: U.S. 60/195,852
; PRIOR FILING DATE: 2000-04-06
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 10
; LENGTH: 21
; TYPE: RNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Synthetic RNA
US-09-828-034-10

Query Match      1.6%; Score 21; DB 9; Length 21;
Best Local Similarity 100.0%; Pred. No. 13;
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGC 76
DB 21 GCGCGCGCGCGCGCGCGCGC 1

RESULT 9
US-10-157-382-7/c
; Sequence 7, Application US/10157382
; Publication No. US20030082668A1
; GENERAL INFORMATION:
; APPLICANT: TAWAI, Katsuyuki
; APPLICANT: MIYAZAKI, Toshiaki
; APPLICANT: WADA, Emi
; APPLICANT: TATSUZAWA, Ayumi
; TITLE OF INVENTION: METHOD FOR MEASURING THE ACTIVITY OF DEACETYLASE
; TITLE OF INVENTION: AND METHOD OF SCREENING FOR INHIBITORS AND ACCELERATORS
; FILE REFERENCE: M3-109PCT-US(CIP)
; CURRENT APPLICATION NUMBER: US/10/157,382
; CURRENT FILING DATE: 2002-05-29
; PRIOR APPLICATION NUMBER: PCT/JP00/08417
; PRIOR FILING DATE: 2000-11-21
; PRIOR APPLICATION NUMBER: JP 1999-338565
; PRIOR FILING DATE: 1999-11-29
; NUMBER OF SEQ ID NOS: 12
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 7
; LENGTH: 39
; TYPE: DNA
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Artificially
; OTHER INFORMATION: Synthesized Sequence
US-10-157-382-7

Query Match      1.6%; Score 21; DB 15; Length 39;
Best Local Similarity 100.0%; Pred. No. 12;
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 560 GTGATGATGCTGATGCTGCTG 580
DB 35 GTGATGATGCTGATGCTGCTG 15
```



RESULT 10  
US-10-157-382-8  
; Sequence 8, Application US/10157382  
; Publication No. US20030082668A1  
; GENERAL INFORMATION:  
; APPLICANT: TAMAI, Katsuyuki  
; APPLICANT: WADA, EMI  
; APPLICANT: MIYAZAKI, Toshiaki  
; APPLICANT: TATSUZAWA, Ayumi  
; TITLE OF INVENTION: METHOD FOR MEASURING THE ACTIVITY OF DEACETYLASE  
; TITLE OF INVENTION: AND METHOD OF SCREENING FOR INHIBITORS AND ACCELERATORS  
; TITLE OF INVENTION: OF THE ENZYME  
; FILE REFERENCE: W3-109PCT-US(CIP)  
; CURRENT APPLICATION NUMBER: US/10/157,382  
; CURRENT FILING DATE: 2002-05-29  
; PRIOR APPLICATION NUMBER: PCT/JP00/08417  
; PRIOR FILING DATE: 2000-11-21  
; PRIOR APPLICATION NUMBER: JP 1999-338565  
; PRIOR FILING DATE: 1999-11-29  
; NUMBER OF SEQ ID NOS: 12  
; SOFTWARE: PatentIn Ver. 2.0  
; SEQ ID NO 8  
; LENGTH: 39  
; TYPE: DNA  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Description of Artificial Sequence:Artificially  
; OTHER INFORMATION: Synthesized Sequence  
US-10-157-382-8  
Query Match 1.6%; Score 21; DB 15; Length 39;  
Best Local Similarity 100.0%; Pred. No. 12;  
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 560 GTGATGATGGTGGTGGTG 580  
DB 9 GTGATGATGGTGGTGGTG 29  
RESULT 11  
US-09-888-326-410  
; Sequence 410, Application US/09888326  
; Publication No. US20030086801A1  
; GENERAL INFORMATION:  
; APPLICANT: Weiner, George  
; APPLICANT: Hartmann, Gunther  
; TITLE OF INVENTION: Methods for Enhancing Antibody-Induced  
; TITLE OF INVENTION: Cell Lysis and Treating Cancer  
; FILE REFERENCE: C1039/7052 (AKS)  
; CURRENT APPLICATION NUMBER: US/09/888,326  
; CURRENT FILING DATE: 2001-06-22  
; PRIOR APPLICATION NUMBER: US 60/213,346  
; PRIOR FILING DATE: 2000-06-22  
; NUMBER OF SEQ ID NOS: 848  
; SOFTWARE: FastSeq for Windows Version 3.0  
; SEQ ID NO 410  
; LENGTH: 20  
; TYPE: DNA  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Synthetic oligonucleotide  
; NAME/KEY: misc feature  
; LOCATION: (0)...(0)  
; OTHER INFORMATION: phosphodiester backbone  
US-09-888-326-410  
Query Match 1.5%; Score 20; DB 10; Length 20;  
Best Local Similarity 100.0%; Pred. No. 41;  
Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 56 GCGCGCGCGCGCGCGCGG 75  
DB 1 GCGCGCGCGCGCGCGCGG 20

RESULT 12  
US-09-776-479-243  
; Sequence 243, Application US/09776479  
; Publication No. US20030087848A1  
; GENERAL INFORMATION:  
; APPLICANT: Bratzler, Robert L.  
; APPLICANT: Fouron, Yves  
; APPLICANT: Bratzler, Robert L.  
; APPLICANT: Fouron, Yves  
; TITLE OF INVENTION: Immunostimulatory Nucleic Acids for the  
; TITLE OF INVENTION: Treatment of Asthma and Allergy  
; FILE REFERENCE: C1037/7013 (HCL/MAT)  
; CURRENT APPLICATION NUMBER: US/09/776,479  
; CURRENT FILING DATE: 2001-02-02  
; PRIOR APPLICATION NUMBER: US 60/179,991  
; PRIOR FILING DATE: 2000-02-03  
; NUMBER OF SEQ ID NOS: 1093  
; SOFTWARE: FastSeq for Windows Version 3.0  
; SEQ ID NO 243  
; LENGTH: 20  
; TYPE: DNA  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Synthetic Sequence  
US-09-776-479-243  
Query Match 1.5%; Score 20; DB 10; Length 20;  
Best Local Similarity 100.0%; Pred. No. 41;  
Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 56 GCGCGCGCGCGCGCGCGG 75  
DB 1 GCGCGCGCGCGCGCGCGG 20  
RESULT 13  
US-10-314-578-243  
; Sequence 243, Application US/10314578  
; Publication No. US20030212026A1  
; GENERAL INFORMATION:  
; APPLICANT: Schetter, Christian  
; APPLICANT: Vollmer, Jorg  
; TITLE OF INVENTION: Immunostimulatory Nucleic Acids  
; FILE REFERENCE: C1039/7035 (HCL/MAT)  
; CURRENT APPLICATION NUMBER: US/10/314,578  
; CURRENT FILING DATE: 2002-12-09  
; PRIOR APPLICATION NUMBER: US 60/156,113  
; PRIOR FILING DATE: 1999-09-25  
; PRIOR APPLICATION NUMBER: US 60/156,135  
; PRIOR FILING DATE: 1999-09-27  
; PRIOR APPLICATION NUMBER: US 60/227,436  
; PRIOR FILING DATE: 2000-08-23  
; NUMBER OF SEQ ID NOS: 1145  
; SOFTWARE: FastSeq for Windows Version 3.0  
; SEQ ID NO 243  
; LENGTH: 20  
; TYPE: DNA  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Synthetic Sequence  
US-10-314-578-243  
Query Match 1.5%; Score 20; DB 13; Length 20;  
Best Local Similarity 100.0%; Pred. No. 41;  
Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 56 GCGCGCGCGCGCGCGCGG 75  
DB 1 GCGCGCGCGCGCGCGCGG 20

RESULT 14

US-09-776-479-243  
 ; Sequence 243, Application US/09776479  
 ; Publication No. US20040067902A9  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Bratzler, Robert L.  
 ; APPLICANT: Petersen, Deanna M.  
 ; APPLICANT: Fouron, Yves  
 ; TITLE OF INVENTION: Immunostimulatory Nucleic Acids for the  
 ; FILE REFERENCE: C1037/7013 (HCL/MAT)  
 ; CURRENT APPLICATION NUMBER: US/09/776,479  
 ; CURRENT FILING DATE: 2001-02-02  
 ; PRIOR APPLICATION NUMBER: US 60/179,991  
 ; PRIOR FILING DATE: 2000-02-03  
 ; NUMBER OF SEQ ID NOS: 1093  
 ; SOFTWARE: FastSeq for Windows Version 3.0  
 ; SEQ ID NO 243  
 ; LENGTH: 20  
 ; TYPE: DNA  
 ; ORGANISM: Artificial Sequence  
 ; FEATURE:  
 ; OTHER INFORMATION: Synthetic Sequence  
 US-09-776-479-243

Query Match 1.5%; Score 20; DB 13; Length 20;  
 Best Local Similarity 100.0%; Pred. No. 41;  
 Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCG 75  
 DB 1 GCGCGCGCGCGCGCGCG 20

RESULT 15

US-10-112-653-235  
 ; Sequence 235, Application US/10112653  
 ; Publication No. US20030050268A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Krieg, Arthur M.  
 ; APPLICANT: Berg, Daniel J.  
 ; TITLE OF INVENTION: IMMUNOSTIMULATORY NUCLEIC ACID FOR  
 ; FILE REFERENCE: C01039/70060 (AWS)  
 ; CURRENT APPLICATION NUMBER: US/10/112,653  
 ; CURRENT FILING DATE: 2002-03-29  
 ; PRIOR APPLICATION NUMBER: US 60/279,642  
 ; PRIOR FILING DATE: 2001-03-29  
 ; NUMBER OF SEQ ID NOS: 1040  
 ; SOFTWARE: FastSeq for Windows Version 3.0  
 ; SEQ ID NO 235  
 ; LENGTH: 20  
 ; TYPE: DNA  
 ; ORGANISM: Artificial Sequence  
 ; FEATURE:  
 ; OTHER INFORMATION: Synthetic Oligonucleotide  
 US-10-112-653-235

Query Match 1.5%; Score 20; DB 15; Length 20;  
 Best Local Similarity 100.0%; Pred. No. 41;  
 Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCG 75  
 DB 1 GCGCGCGCGCGCGCGCG 20

Search completed: July 1, 2004, 23:29:54  
 Job time : 648 secs

GenCore version 5.1.6  
Copyright (c) 1993 - 2004 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: July 1, 2004, 19:03:10 ; Search time 3776 Seconds  
(without alignments)  
10447.018 Million cell updates/sec

Title: US-09-934-249-1  
Perfect score: 1321  
Sequence: 1 cgaccgcgtctggagcga.....ctgcgtaggtagaaaggcag 1321

Scoring table: OLIGO NUC  
Gapop 60.0, Gapext 60.0

Searched: 27513289 seqs, 14931090276 residues

Word size : 0

Total number of hits satisfying chosen parameters: 138346

Minimum DB seq length: 0

Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database :

- EST.\*
- 1: em\_estba.\*
  - 2: em\_esthum.\*
  - 3: em\_estin.\*
  - 4: em\_estmu.\*
  - 5: em\_estov.\*
  - 6: em\_estpl.\*
  - 7: em\_estro.\*
  - 8: em\_hic.\*
  - 9: gb\_est1.\*
  - 10: gb\_est2.\*
  - 11: gb\_est3.\*
  - 12: gb\_est4.\*
  - 13: gb\_est5.\*
  - 14: gb\_est6.\*
  - 15: em\_estfun.\*
  - 16: em\_estom.\*
  - 17: em\_gss\_hum.\*
  - 18: em\_gss\_inv.\*
  - 19: em\_gss\_pln.\*
  - 20: em\_gss\_vrt.\*
  - 21: em\_gss\_fun.\*
  - 22: em\_gss\_man.\*
  - 23: em\_gss\_mus.\*
  - 24: em\_gss\_pro.\*
  - 25: em\_gss\_rtd.\*
  - 26: em\_gss\_pg.\*
  - 27: em\_gss\_vrl.\*
  - 28: gb\_gss1.\*
  - 29: gb\_gss2.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

# SUMMARIES

| Result No. | Score | Query Match | Length | ID | Description |
|------------|-------|-------------|--------|----|-------------|
| 1          | 23    | 1.7         | 33     | 10 | BE886705    |
| 2          | 22    | 1.7         | 24     | 14 | CF291636    |
| 3          | 22    | 1.7         | 50     | 9  | AU107980    |
| 4          | 20    | 1.5         | 35     | 2  | HS000944    |

|    |    |     |    |    |           |
|----|----|-----|----|----|-----------|
| 5  | 20 | 1.5 | 39 | 10 | BE8871689 |
| 6  | 20 | 1.5 | 48 | 29 | CG779308  |
| 7  | 19 | 1.4 | 39 | 10 | BE8871689 |
| 8  | 18 | 1.4 | 24 | 14 | CF291636  |
| 9  | 18 | 1.4 | 33 | 10 | BE886705  |
| 10 | 18 | 1.4 | 34 | 28 | AZ761910  |
| 11 | 18 | 1.4 | 43 | 28 | CC199634  |
| 12 | 18 | 1.4 | 46 | 28 | AZ993993  |
| 13 | 18 | 1.4 | 48 | 29 | CG779308  |
| 14 | 18 | 1.4 | 50 | 9  | AU107980  |
| 15 | 17 | 1.3 | 27 | 28 | AZ842796  |
| 16 | 17 | 1.3 | 33 | 28 | AZ401045  |
| 17 | 17 | 1.3 | 35 | 2  | HS000944  |
| 18 | 17 | 1.3 | 41 | 28 | AZ779226  |
| 19 | 17 | 1.3 | 43 | 28 | CC199634  |
| 20 | 17 | 1.3 | 44 | 12 | BI767274  |
| 21 | 17 | 1.3 | 44 | 28 | AZ635754  |
| 22 | 17 | 1.3 | 45 | 12 | BI772215  |
| 23 | 17 | 1.3 | 45 | 12 | BI772215  |
| 24 | 17 | 1.3 | 49 | 10 | BF970690  |
| 25 | 17 | 1.3 | 50 | 9  | AU104810  |
| 26 | 16 | 1.2 | 20 | 14 | CF317946  |
| 27 | 16 | 1.2 | 20 | 14 | CF317946  |
| 28 | 16 | 1.2 | 37 | 14 | CF328207  |
| 29 | 16 | 1.2 | 40 | 10 | BE397229  |
| 30 | 16 | 1.2 | 41 | 10 | BE264159  |
| 31 | 16 | 1.2 | 44 | 12 | BI767274  |
| 32 | 16 | 1.2 | 50 | 9  | AU102484  |
| 33 | 16 | 1.2 | 50 | 9  | AU102609  |
| 34 | 16 | 1.2 | 50 | 9  | AU102610  |
| 35 | 16 | 1.2 | 50 | 9  | AU103688  |
| 36 | 16 | 1.2 | 50 | 9  | AU103691  |
| 37 | 16 | 1.2 | 50 | 9  | AU104498  |
| 38 | 16 | 1.2 | 50 | 9  | AU104499  |
| 39 | 16 | 1.2 | 50 | 9  | AU104500  |
| 40 | 16 | 1.2 | 50 | 9  | AU104501  |
| 41 | 16 | 1.2 | 50 | 9  | AU104502  |
| 42 | 16 | 1.2 | 50 | 9  | AU104515  |
| 43 | 16 | 1.2 | 50 | 9  | AU104520  |
| 44 | 16 | 1.2 | 50 | 9  | AU104811  |
| 45 | 16 | 1.2 | 50 | 9  | AU106823  |

## ALIGNMENTS

RESULT 1  
BE886705  
LOCUS  
DEFINITION  
601507961F1 NIH\_MGC\_71 Homo sapiens cDNA clone IMAGE:3909591 5', linear EST 20-OCT-2000  
mRNA sequence.  
ACCESSION  
BE886705  
VERSION  
BE886705.1 GI:10341256  
KEYWORDS  
EST.  
SOURCE  
Homo sapiens (human)  
ORGANISM  
Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Homidae; Homo.  
REFERENCE  
1 (bases 1 to 33)  
NIH-MGC <http://mgc.nci.nih.gov/>.  
AUTHORS  
National Institutes of Health, Mammalian Gene Collection (MGC)  
TITLE  
Unpublished (1999)  
JOURNAL  
COMMENT  
Contact: Robert Strausberg, Ph.D.  
Email: cgapbs-r@mail.nih.gov  
Tissue Procurement: ATCC  
CDNA Library Preparation: Life Technologies, Inc.  
CDNA Library Arrayed by: The I.M.A.G.E. Consortium (LMNL)  
DNA Sequencing by: Incyte Genomics, Inc.  
Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at: <http://image.llnl.gov>  
Plate: L1AM9723 row: i column: 16  
High quality sequence stop: 33.

FEATURES  
sourceLocation/Qualifiers  
1. .33

/organism="Homo sapiens"  
/mol\_type="mRNA"  
/db\_xref="taxon:9606"  
/clone="IMAGE:3909591"  
/tissue\_type="leiomyosarcoma"  
/lab\_host="DH10B (phage-resistant)"  
/clone\_lib="NH1\_MGC\_71"  
/note="Organ: uterus; Vector: pCMV-SPORT6; Site\_1: NotI;  
Site\_2: SalI; Cloned unidirectionally. Primer: Oligo dt.  
Average insert size 2.1 kb."

## ORIGIN

Query Match 1.7%; Score 23; DB 10; Length 33;  
Best Local Similarity 100.0%; Pred. No. 5.6e+02;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 54 GAGCGCGCGCGCGCGCGCGCG 76  
DB 11 GAGCGCGCGCGCGCGCGCGCG 33

## RESULT 2

## CF291636

## LOCUS

DEFINITION 14ROOT--02-C09.g1 Rice root plasmid cDNA library (14ROOT) Oryza  
sativa cDNA clone 14ROOT--02-C09, mRNA sequence.

## ACCESSION

## CF291636

## VERSION

## CF291636.1

## KEYWORDS

## EST.

## SOURCE

## Oryza sativa

## ORGANISM

Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;  
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;  
Ehrhartoideae; Oryzaceae; Oryza.

## REFERENCE

## AUTHORS

## TITLE

## JOURNAL

## COMMENT

## FEATURES

## source

Location/Qualifiers  
1. .24

/organism="Oryza sativa"  
/mol\_type="mRNA"  
/cultivar="Nackdong"  
/db\_xref="taxon:4530"  
/clone="14ROOT--02-C09"  
/tissue\_type="root"  
/dev\_stage="14 days after germination"  
/lab\_host="E.coli DH10B"  
/clone\_lib="Rice root plasmid cDNA library (14ROOT)"  
/note="Vector: pCR4-TOPO; Site\_1: EcoRI; mRNA was capped  
with oligoribonucleotides and then used as templates for  
RT-PCR."

## ORIGIN

Query Match 1.7%; Score 22; DB 14; Length 24;  
Best Local Similarity 100.0%; Pred. No. 1.4e+03;  
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCG 77

DB 3 GCGCGCGCGCGCGCGCGCGCG 24

## RESULT 3

## AUI07980

## LOCUS

## DEFINITION

## AUI07980

## ACCESSION

## AUI07980

## VERSION

## AUI07980.1

## KEYWORDS

## EST.

## SOURCE

## Homo sapiens (human)

## ORGANISM

Eukaryota; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

## REFERENCE

## AUTHORS

## TITLE

## JOURNAL

## MEDLINE

## PUBMED

## COMMENT

## Contact: Yutaka Suzuki

## Department of Virology

## Institute of Medical Science, University of Tokyo

## 4-6-1, Shirokanedai, Minatoku, Tokyo 108-8639, Japan

## Email: ysuzuki@ims.u-tokyo.ac.jp

## Suzuki, Y., Yoshitomo-Nakagawa, K., Maruyama, K., Suyama, A. and

## Sugano, S. Construction and characterization of a full

## length-enriched and a 5'-end-enriched cDNA library. Gene 200 (1-2),

## 149-156 (1997).

## 149-156 (1997).

## 149-156 (1997).

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## 149-156 (1997).

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## 149-156 (1997).

## 149-156 (1997).

## 149-156 (1997).

## 149-156 (1997).

## 149-156 (1997).

```

XX CC Clone from S. Wiemann, sequenced by AGOWA within the cDNA
CC sequencing consortium of the German Genome Project
CC si sequence also available
CC This clone is available at the RZPD in Berlin
CC Please contact the RZPD: Ressourcenzentrum, Heubnerweg 6, 14059
CC Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de
XX FH Key Location/Qualifiers
FH source 1. .35
FT /db_xref="taxon:9606"
FT /mol_type="rRNA"
FT /organism="Homo sapiens"
FT /clone="DKFZp434L174"
FT /clone_lib="434 (synonym: htes3). Vector pSport1; host
FT DH10B; sites NotI + SalI"
FT /dev_stage="adult"
FT /tissue_type="testis"
XX SQ Sequence 35 BP; 1 A; 12 C; 20 G; 2 T; 0 other;
Query Match 1.5%; Score 20; DB 2; Length 35;
Best Local Similarity 100.0%; Pred. No. 7.3e+03;
Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 53 GGAGCGCGCGCGCGCGCG 72
Db 9 GGAGCGCGCGCGCGCGCG 28

RESULT 5
BE871689
LOCUS BE871689 39 bp mRNA linear EST 20-OCT-2000
DEFINITION 601449550F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3853381 5',
mRNA sequence.
ACCESSION BE871689
VERSION BE871689.1 GI:10320465
KEYWORDS EST.
SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE 1 (bases 1 to 39)
AUTHORS NIH-MGC http://mgc.ncl.nih.gov/
TITLE National Institutes of Health, Mammalian Gene Collection (MGC)
JOURNAL Unpublished (1999)
COMMENT Contact: Robert Strausberg, Ph.D.
Email: cgapbs@mail.nih.gov
Tissue Procurement: ATCC
CDNA Library Preparation: Life Technologies, Inc.
CDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Incyte Genomics, Inc.
Clone distribution: MGC clone distribution information can be
found through the I.M.A.G.E. Consortium/LLNL at:
http://image.llnl.gov
Plate: LLAM9577 row: c column: 14
High quality sequence stop: 39.
FEATURES
source 1. .35
Location/Qualifiers
/organism="Homo sapiens"
/mol_type="mRNA"
/db_xref="taxon:9606"
/clone="IMAGE:3853381"
/tissue_type="adenocarcinoma"
/lab_host="DH10B (phage-resistant)"
/clone_lib="NIH_MGC_65"
/Notes="Organ: colon; Vector: pCMV-Sport6; Site:1: NotI;
Site:2: SalI; Cloned unidirectionally. Primer: Oligo dr.
Average insert size 1.8 kb. Library constructed by Life
Technologies."
ORIGIN

```

```

Query Match 1.5%; Score 20; DB 10; Length 39;
Best Local Similarity 100.0%; Pred. No. 7.3e+03;
Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GGCGCGCGCGCGCGCGCG 75
Db 11 GGCGCGCGCGCGCGCGCG 30

RESULT 6
CG779308
LOCUS CG779308 48 bp DNA linear GSS 29-OCT-2003
DEFINITION 1123033B10.2EL x1 1123 - RescueMu Grid L Zea mays genomic
survey sequence.
ACCESSION CG779308
VERSION CG779308.1 GI:38040097
KEYWORDS GSS.
SOURCE Zea mays
ORGANISM Zea mays
Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; PACCAD
clade; Panicoideae; Andropogoneae; Zea.
REFERENCE 1 (bases 1 to 48)
AUTHORS Walbot,V.
TITLE Maize genomic sequences found using engineered RescueMu transposon
JOURNAL Unpublished (2001)
COMMENT Contact: Walbot V
Department of Biological Sciences
Stanford University
855 California Ave, Palo Alto, CA 94304, USA
Tel: 650 723 2227
Fax: 650 725 8221
Email: walbot@stanford.edu
Possible ligation site of ends cut by 2 different endonucleases.
Reverse complemented post-ligation sequence from source sequence.
Plate: 1123033 row: 14
Class: transposon-tagged.
FEATURES
source 1. .48
Location/Qualifiers
/organism="Zea mays"
/mol_type="genomic DNA"
/cultivar="mixed background W23/A188/B73/K55"
/db_xref="taxon:4577"
/tissue_type="leaf"
/dev_stage="adult"
/lab_host="DH10B"
/clone_lib="1123 - RescueMu Grid L"
/Notes="Organ: leaf; Vector: RescueMu (engineered from
pBlueScript backbone); Site 1: BamHI; Site 2: BglII;
RescueMu is a 4.9 kb, modified maize Mu transposon
designed to allow plasmid rescue from total genomic DNA.
Mu elements insert preferentially into transcription
units. For more information on RescueMu, go to the web
site 'www.zmdb.iastate.edu' and follow the links for
'RescueMu.' Grid L was grown in Molokai in 2001. DNA was
extracted from leaf strips, double digested using BamHI
and BglII, and ligated to form circular plasmids. DH10B
cells were transformed and then screened on LB plates with
ampicillin."
ORIGIN
Query Match 1.5%; Score 20; DB 29; Length 48;
Best Local Similarity 100.0%; Pred. No. 7.2e+03;
Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 57 GCGCGCGCGCGCGCGCGCG 76
Db 8 GCGCGCGCGCGCGCGCGCG 27

RESULT 7
BE871689/c
LOCUS BE871689 39 bp mRNA linear EST 20-OCT-2000

```

```

DEFINITION 601449550F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3853381 5',
            mRNA sequence.
ACCESSION  BB871689
VERSION     BB871689.1 GI:10320465
KEYWORDS   EST.
SOURCE      Homo sapiens (human)
ORGANISM    Homo sapiens
            Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE   1 (bases 1 to 39)
            NIH-MGC http://mgc.nci.nih.gov/.
            National Institutes of Health, Mammalian Gene Collection (MGC)
            Unpublished (1999)
AUTHORS     Contact: Robert Strausberg, Ph.D.
TITLE       Tissue Procurement: ATCC
JOURNAL     CNA Library Preparation: Life Technologies, Inc.
COMMENT     DNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
            DNA sequencing by: Incyte Genomics, Inc.
            Clone distribution: MGC clone distribution information can be
            found through the I.M.A.G.E. Consortium/LLNL at:
            http://image.llnl.gov
            Plate: LAM9577 row: C column: 14
            High quality sequence stop: 39.
FEATURES    Location/Qualifiers
            1..39
            /organism="Homo sapiens"
            /mol_type="mRNA"
            /db_xref="taxon:9606"
            /clone="IMAGE:3853381"
            /tissue_type="adenocarcinoma"
            /lab_host="DH10B (phage-resistant)"
            /clone_lib="NIH MGC_65"
            /note="Organ: Colon; Vector: pCMV-SPORT6; Site 1: NotI;
            Site 2: SalI; Cloned unidirectionally. Primer: Oligo dt.
            Average insert size 1.8 kb. Library constructed by Life
            Technologies."
ORIGIN
Query Match 1.4%; Score 19; DB 10; Length 39;
Best Local Similarity 100.0%; Pred. No. 1.7e+04;
Matches 19; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 440 ACCGCGCGCGCGCGCGCGCG 458
    |||||
DB 31 ACCGCGCGCGCGCGCGCGCG 13

RESULT 8
CF291636/24 bp mRNA linear EST 14-AUG-2003
LOCUS 14800T-02-C09.g1 Rice root plasmid cDNA library (14800T) Oryza
DEFINITION sativa cDNA clone 14800T-02-C09, mRNA sequence.
ACCESSION  CF291636
VERSION     CF291636.1 GI:33660669
KEYWORDS   EST.
SOURCE      Oryza sativa
            Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
            Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
            Ehrhacidae; Oryzaceae; Oryza.
            1 (bases 1 to 24)
            Kim, J.S., Jun, K.M., Cheong, P.J., Kim, M.J., Lee, T.H., Shin, Y.C.,
            Song, S.I., Kim, J.K., Kim, Y.-K. and Nahm, B.H.
            Large-scale Sequencing Analysis of Rice ESTs
            Unpublished (2003)
AUTHORS     Contact: Nahm B.H.
TITLE       Genomics and Genetics Institute, GreenGene Biotech Inc.; Division
JOURNAL     of Bioscience and Bioinformatics, Myongui University
COMMENT     Yongsin, Kyeonggi, Korea
            Tel: 82 31 330 6193
            Fax: 82 31 321 6355
            Email: bhnahm@gbio.com, bhnahm@bio.myongji.ac.kr.

```

```

FEATURES    Location/Qualifiers
            1..24
            /organism="Oryza sativa"
            /mol_type="mRNA"
            /cultivar="Nackdong"
            /db_xref="taxon:4530"
            /clone="14800T-02-C09"
            /tissue_type="root"
            /lab_host="E.coli DH10B"
            /note="Rice root plasmid cDNA library (14800T)"
            /note="Vector: PCR4-TOPO; Site 1: EcoRI; mRNA was capped
            with oligoribonucleotides and then used as templates for
            RT-PCR."
ORIGIN
Query Match 1.4%; Score 18; DB 14; Length 24;
Best Local Similarity 100.0%; Pred. No. 4.2e+04;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 441 CGCGCGCGCGCGCGCGCG 458
    |||||
DB 22 CGCGCGCGCGCGCGCGCG 5

RESULT 9
BE886705/33 bp mRNA linear EST 20-OCT-2000
LOCUS 601507961F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909591 5',
DEFINITION tRNA sequence.
ACCESSION  BE886705
VERSION     BE886705.1 GI:10341256
KEYWORDS   EST.
SOURCE      Homo sapiens (human)
ORGANISM    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
            1 (bases 1 to 33)
            NIH-MGC http://mgc.nci.nih.gov/.
            National Institutes of Health, Mammalian Gene Collection (MGC)
            Unpublished (1999)
            Contact: Robert Strausberg, Ph.D.
            Email: cgabbs-remail.nih.gov
            Tissue Procurement: ATCC
            CNA Library Preparation: Life Technologies, Inc.
            DNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
            DNA sequencing by: Incyte Genomics, Inc.
            Clone distribution: MGC clone distribution information can be
            found through the I.M.A.G.E. Consortium/LLNL at:
            http://image.llnl.gov
            Plate: LAM9723 row: i column: 16
            High quality sequence stop: 33.
FEATURES    Location/Qualifiers
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            /organism="Homo sapiens"
            /mol_type="mRNA"
            /db_xref="taxon:9606"
            /clone="IMAGE:3909591"
            /tissue_type="leiomyosarcoma"
            /lab_host="DH10B (phage-resistant)"
            /clone_lib="NIH MGC_71"
            /note="Organ: uterus; Vector: pCMV-SPORT6; Site 1: NotI;
            Site 2: SalI; Cloned unidirectionally. Primer: Oligo dt.
            Average insert size 2.1 kb."
ORIGIN
Query Match 1.4%; Score 18; DB 10; Length 33;
Best Local Similarity 100.0%; Pred. No. 4.1e+04;
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 441 CGCGCGCGCGCGCGCGCG 458
    |||||
DB 32 CGCGCGCGCGCGCGCGCG 15

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RESULT 10
LOCUS      AZ761910
DEFINITION IM0556C05R Mouse 10kb plasmid UUGC1M library Mus musculus genomic
            clone UUGC1M0556C05 R, genomic survey sequence.
ACCESSION  AZ761910
VERSION     AZ761910.1 GI:12871328
KEYWORDS    GSS.
SOURCE      Mus musculus (house mouse)
ORGANISM    Mus musculus
            Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
REFERENCE   1 (bases 1 to 34)
AUTHORS     Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
            Reilly,M., Rose,M., Rose,R., Stokes,R., Tingey,A., von
            Niederhausern,A. and Wright,D.,Weiss,R.
TITLE       Mouse whole genome scaffolding with paired end reads from 10kb
            plasmid inserts
JOURNAL     Unpublished (2000)
COMMENT     Contact: Robert B. Weiss
            University of Utah
            Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT
            84112, USA
            Tel: 801 585 5606
            Fax: 801 585 7177
            Email: ddunn@genetics.utah.edu
            Insert Length: 10000 Std Error: 0.00
            Plate: 0556 row: C column: 05
            Seq primer: CACACAGGAAACAGCTATGACC
            Class: plasmid ends
            High quality sequence stop: 34.
FEATURES   source
            1..34
                Location/Qualifiers
                1..34
                    /organism="Mus musculus"
                    /mol_type="genomic DNA"
                    /strain="C57BL/6J"
                    /db_xref="taxon:10090"
                    /clone="UUGC1M0556C05"
                    /sex="Male"
                    /lab_host="E. Coli strain XL10-Gold, T1-resistant, F-"
                    /clone_lib="Mouse 10kb plasmid UUGC1M library"
                    /note="Vector: PWD42nv; Purified genomic DNA from M.
                    musculus C57BL/6J (male) was obtained from the Jackson
                    Laboratory Mouse DNA Resource
                    (http://www.jax.org/resources/documents/dnares/). The DNA
                    was hydrodynamically sheared by repeated passage through a
                    0.005 inch orifice at constant velocity. The sheared DNA
                    was blunt end-repaired with T4 DNA polymerase and T4
                    polynucleotide kinase. Adaptor oligonucleotides were
                    ligated to the blunt ends in high molar excess. The
                    adaptor DNA was purified and size-selected for a 9.5 to
                    10.5 kb range using preparative agarose gel
                    electrophoresis. Vector DNA was prepared from a derivative
                    of PWD42 (gi|4732114|gb|AF129072.1), a copy-number
                    inducible derivative of plasmid R1. The vector was ligated
                    with adaptors complementary to the insert adaptors and
                    purified. The sheared, adaptor mouse DNA was annealed to
                    adaptor vector DNA, and transformed into
                    chemically-competent E. coli XL10-Gold (Stratagene) cells
                    and selected for ampicillin resistance."
ORIGIN
            Query Match      1.4%; Score 18; DB 28; Length 34;
            Best Local Similarity 100.0%; Pred. No. 4.1e+04;
            Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY      554 GTGGTGGTGTGATGTGTG 571
Db      14 GTGGTGGTGTGATGTGTG 31

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RESULT 11
LOCUS      CC199634
DEFINITION XH740 BayGenomics Gene Trap Library pGTILxf Mus musculus cDNA, mRNA
            sequence.
ACCESSION  CC199634
VERSION     CC199634.1 GI:30479674
KEYWORDS    GSS.
SOURCE      Mus musculus (house mouse)
ORGANISM    Mus musculus
            Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
REFERENCE   1 (bases 1 to 43)
AUTHORS     BayGenomics.
            Email: info@baygenomics.ucsf.edu
TITLE       Sequence tag generated by 5' RAGE of total RNA from gene trap ES
            cell line. ES cell lines harboring insertion mutation of target
            gene are available upon request from BayGenomics. Annotation
            information available from
            http://baygenomics.ucsf.edu/cgi-bin/BaySearch.py?OPTION=EXACT&TYPE=
            CELL_LINE&KEY=XH740
            Class: Gene Trap.
FEATURES   Location/Qualifiers
            1..43
                /organism="Mus musculus"
                /mol_type="mRNA"
                /strain="129 OLA"
                /db_xref="taxon:10090"
                /sex="Male"
                /cell_type="Embryonic stem cell"
                /clone_lib="BayGenomics Gene Trap Library pGTILxf"
                /note="Vector: pGTILxf"
ORIGIN
            Query Match      1.4%; Score 18; DB 28; Length 43;
            Best Local Similarity 100.0%; Pred. No. 4e+04;
            Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY      56 GGCGCGCGCGCGCGCGCGCGC 73
Db      25 GGCGCGCGCGCGCGCGCGCGC 42

RESULT 12
LOCUS      AZ993993
DEFINITION 2M0279E13F Mouse 10kb plasmid UUGC2M library Mus musculus genomic
            clone UUGC2M0279E13 F, genomic survey sequence.
ACCESSION  AZ993993
VERSION     AZ993993.1 GI:13865220
KEYWORDS    GSS.
SOURCE      Mus musculus (house mouse)
ORGANISM    Mus musculus
            Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
REFERENCE   1 (bases 1 to 46)
AUTHORS     Dunn,D., Aoyagi,A., Barber,M., Beacorn,T., Duval,B., Hamil,C.,
            Reilly,M., Rose,M., Rose,R., Stokes,R., Tingey,A., von
            Niederhausern,A. and Wright,D.,Weiss,R.
TITLE       Mouse whole genome scaffolding with paired end reads from 10kb
            plasmid inserts
JOURNAL     Unpublished (2000)
COMMENT     Contact: Robert B. Weiss
            University of Utah
            Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT

```

84112, USA  
 Tel: 801 585 5606  
 Fax: 801 585 7177  
 Email: ddunn@genetics.utah.edu  
 Insert Length: 10000 Std Error: 0.00  
 Plate: 0279 row: E column: 13  
 Seq primer: CGTGTAAACGACGCGCAGT  
 Class: Plasmid ends  
 High quality sequence stop: 46.

## FEATURES

source

1. .46  
 /organism="Mus musculus"  
 /mol\_type="genomic DNA"  
 /strain="C57BL/6J"  
 /db\_xref="taxon:10090"  
 /clone="UGC2M0279E13"  
 /sex="female"  
 /lab\_host="E. coli strain XL10-Gold, T1-resistant, P-"  
 /clone\_lib="Mouse 10kb plasmid UGC2M library"  
 /notes="Vector: PWD42nv; Purified genomic DNA from M. musculus C57BL/6J (female) was obtained from the Jackson Laboratory Mouse DNA Resource  
 (http://www.jax.org/resources/documents/dnares/). The DNA was hydrodynamically sheared by repeated passage through a 0.005 inch orifice at constant velocity. The sheared DNA was blunt end-repaired with T4 DNA polymerase and T4 polynucleotide kinase. Adaptor oligonucleotides were ligated to the blunt ends in high molar excess. The adaptor DNA was purified and size-selected for a 9.5 to 10.5 kb range using preparative agarose gel electrophoresis. Vector DNA was prepared from a derivative of PWD42 [gi|4732114|gb|AF129072.1], a copy-number inducible derivative of plasmid R1. The vector was ligated with adaptors complementary to the insert adaptors and purified. The sheared, adaptor mouse DNA was annealed to adaptor vector DNA, and transformed into chemically-competent E. coli XL10-Gold (Stratagene) cells and selected for ampicillin resistance."

## ORIGIN

Query Match 1.4%; Score 18; DB 28; Length 46;  
 Best Local Similarity 100.0%; Pred. No. 4e+04;  
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 561 TCATCATGCTGATGCTGC 578

Db 29 TCATCATGCTGATGCTGC 46

## RESULT 13

CG779308/c

LOCUS

DEFINITION 1123033B10.2BLx1 1123 - RescueMu Grid L Zea mays genomic, Genomic survey sequence.

ACCESSION

CG779308

VERSION

CG779308.1

KEYWORDS

GSS.

SOURCE

Zea mays

ORGANISM

Zea mays

REFERENCE

AUTHORS

TITLE

JOURNAL

COMMENT

CONTACT: Walbot V

Department of Biological Sciences

Stanford University

855 California Ave, Palo Alto, CA 94304, USA

Tel: 650 723 2227

Fax: 650 725 8221

Email: walbot@stanford.edu

Possible ligation site of ends cut by 2 different endonucleases.  
 Reverse complemented post-ligation sequence from source sequence.  
 Plate: 1123033 row: 14  
 Class: transposon-tagged.  
 Location/Qualifiers

## FEATURES

source

1. .48  
 /organism="Zea mays"  
 /mol\_type="genomic DNA"  
 /cultivar="mixed background W23/A188/B73/K55"  
 /db\_xref="taxon:4577"  
 /tissue\_type="leaf"  
 /dev\_stage="adult"  
 /lab\_host="DH10B"  
 /clone\_lib="1123 - RescueMu Grid L"  
 /note="Organ: leaf; Vector: RescueMu (engineered from pBluescript backbone); Site 1: BamHI; Site 2: BglII; RescueMu is a 4.9 kb, modified maize Mu transposon designed to allow plasmid rescue from total genomic DNA. Mu elements insert preferentially into transcription units. For more information on RescueMu, go to the web site 'www.zmdb.iastate.edu' and follow the links for 'RescueMu.' Grid L was grown in Molokai in 2001. DNA was extracted from leaf strips double digested using BamHI and BglII, and ligated to form circular plasmids. DH10B cells were transformed and then screened on LB plates with ampicillin."

## ORIGIN

Query Match 1.4%; Score 18; DB 29; Length 48;  
 Best Local Similarity 100.0%; Pred. No. 4e+04;  
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 441 CCGCCGCGCGCGCGCGCGC 458

Db 26 CCGCCGCGCGCGCGCGCGC 9

## RESULT 14

AU107980/c

LOCUS

DEFINITION

ZRV62238, mRNA sequence.

ACCESSION

AU107980

VERSION

AU107980.1

KEYWORDS

EST.

SOURCE

Homo sapiens (human)

ORGANISM

Homo sapiens

REFERENCE

AUTHORS

Suzuki, Y., Taira, H., Tsunoda, T., Mizushima-Sugano, J., Sese, J., Hata, H., Ota, T., Isogai, T., Tanaka, T., Morishita, S., Okubo, K., Sakaki, Y., Nakamura, Y., Suyama, A. and Sugano, S.

TITLE

Diverse transcriptional initiation revealed by fine, large-scale mapping of mRNA start sites

JOURNAL

EMBO Rep. 2 (5), 388-393 (2001)

MEDLINE

21270072

PUBMED

11375929

COMMENT

Contact: Yutaka Suzuki

Department of Virology

Institute of Medical Science, University of Tokyo

4-6-1, Shirokanedai, Minatoku, Tokyo 108-8639, Japan

Email: yusuzuki@ims.u-tokyo.ac.jp

Suzuki, Y., Yoshitomo-Nakagawa, K., Maruyama, K., Suyama, A. and Sugano, S.

TITLE

Construction and characterization of a full length-enriched and a 5'-end-enriched cDNA library. Gene 200 (1-2), 149-156 (1997).

JOURNAL

149-156 (1997).

COMMENT

Location/Qualifiers

1. .50

/organism="Homo sapiens"

/mol\_type="mRNA"

/db\_xref="taxon:9606"

/clone="ZRV62238"



/clone\_lib="Sugano Homo sapiens cDNA library"

## ORIGIN

Query Match 1.4%; Score 18; DB 9; Length 50;  
Best Local Similarity 100.0%; Pred. No. 4e+04;  
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 441 CGCGCGCGCGCGCGCG 458  
|||||  
Db 29 CGCGCGCGCGCGCGCG 12

## RESULT 15

AZ842796

LOCUS

DEFINITION 27 bp DNA linear GSS 20-FEB-2001  
2M0141120F Mouse 10kb plasmid UUGC1M library Mus musculus genomic  
clone UUGC2M0141120 F, genomic survey sequence.

ACCESSION AZ842796

VERSION AZ842796.1 GI:13012704

KEYWORDS GSS.

SOURCE Mus musculus (house mouse)

## ORGANISM

Mus musculus  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.

## REFERENCE

AUTHORS

## TITLE

Mouse whole genome scaffolding with paired end reads from 10kb  
plasmid inserts

## JOURNAL

COMMENT

Unpublished (2000)

Contact: Robert B. Weiss

University of Utah Genome Center

University of Utah

Rm. 308, Biomedical Polymers Research Bldg., 20 S. 2030 E., SLC, UT

84112, USA

Tel: 801 585 5606

Fax: 801 585 7177

Email: ddunn@genetics.utah.edu

Insert Length: 10000 Std Error: 0.00

Plate: 0141 row: I column: 20

Seq primer: CGTTGTAAACGACGCGCCAGT

Class: plasmid ends

High quality sequence stop: 27.

## FEATURES

source

1. .27  
Location/Qualifiers  
/organism="Mus musculus"  
/mol\_type="genomic DNA"  
/strain="C57BL/6J"  
/db\_xref="taxon:10090"  
/clone="UUGC2M0141120"  
/sex="Male"

/lab\_host="B. Coli strain XL10-Gold, T1-resistant, F-"  
/clone\_lib="Mouse 10kb plasmid UUGC1M library"  
/note="Vector: PWD42nv; Purified genomic DNA from M.  
musculus C57BL/6J (male) was obtained from the Jackson  
Laboratory Mouse DNA Resource  
(http://www.jax.org/resources/documents/dnares/). The DNA  
was hydrodynamically sheared by repeated passage through a  
0.005 inch orifice at constant velocity. The sheared DNA  
was blunt end-repaired with T4 DNA polymerase and T4  
polynucleotide kinase. Adaptor oligonucleotides were  
ligated to the blunt ends in high molar excess. The  
adaptored DNA was purified and size-selected for a 9.5 to  
10.5 kb range using preparative agarose gel  
electrophoresis. Vector DNA was prepared from a derivative  
of PWD42 (GI|4732114|gb|AF129072.1), a copy-number  
inducible derivative of plasmid R1. The vector was ligated  
with adaptors complementary to the insert adaptors and  
purified. The sheared, adaptored mouse DNA was annealed to  
adaptored vector DNA, and transformed into  
chemically-competent E. coli XL10-Gold (Stratagene) cells

and selected for ampicillin resistance."

## ORIGIN

Query Match 1.3%; Score 17; DB 28; Length 27;  
Best Local Similarity 100.0%; Pred. No. 9.7e+04;  
Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 567 TGGTGATGGTGGTGGTG 583  
|||||  
Db 11 TGGTGATGGTGGTGGTG 27

Search completed: July 1, 2004, 21:51:07

Job time : 3778 secs

GenCore version 5.1.6  
Copyright (c) 1993 - 2004 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: July 1, 2004, 19:04:19 ; Search time 115 Seconds  
(without alignments)  
6374.697 Million cell updates/sec

Title: US-09-934-249-1  
Perfect score: 1321  
Sequence: 1 cgaccggctctggagcga.....ctgcgtaggtaaaaggcag 1321

Scoring table: OUIGO.NUC  
Gapop 60.0 , Gapext 60.0

Searched: 682709 seqs, 277475446 residues

Word size : 0

Total number of hits satisfying chosen parameters: 839752

Minimum DB seq length: 0  
Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database : Issued Patents\_NA.\*  
1: /cgm2\_6/ptodata/2/ina/5A.COMB.seq.\*  
2: /cgm2\_6/ptodata/2/ina/5B.COMB.seq.\*  
3: /cgm2\_6/ptodata/2/ina/6A.COMB.seq.\*  
4: /cgm2\_6/ptodata/2/ina/6B.COMB.seq.\*  
5: /cgm2\_6/ptodata/2/ina/PCTUS.COMB.seq.\*  
6: /cgm2\_6/ptodata/2/ina/backfiles1.seq.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

# SUMMARIES

| Result No. | Score | Query Match | Length | ID | Description        |
|------------|-------|-------------|--------|----|--------------------|
| 1          | 24    | 1.8         | 30     | 4  | US-09-475-947A-332 |
| 2          | 24    | 1.8         | 50     | 4  | US-09-930-181-14   |
| 3          | 23    | 1.7         | 25     | 1  | US-08-374-144-3    |
| 4          | 23    | 1.7         | 25     | 1  | US-08-775-164-3    |
| 5          | 23    | 1.7         | 25     | 2  | US-08-775-609-3    |
| 6          | 23    | 1.7         | 25     | 2  | US-08-775-607-3    |
| 7          | 23    | 1.7         | 25     | 5  | PCT-US93-06828-3   |
| 8          | 23    | 1.7         | 30     | 1  | US-08-068-747-2    |
| 9          | 23    | 1.7         | 33     | 1  | US-08-068-747-7    |
| 10         | 23    | 1.7         | 45     | 4  | US-09-497-933A-22  |
| 11         | 23    | 1.7         | 45     | 4  | US-09-497-933A-19  |
| 12         | 22    | 1.7         | 22     | 4  | US-08-769-482-8    |
| 13         | 21    | 1.6         | 21     | 2  | US-08-863-639A-52  |
| 14         | 21    | 1.6         | 21     | 2  | US-08-863-639A-55  |
| 15         | 21    | 1.6         | 21     | 2  | US-08-863-639A-56  |
| 16         | 21    | 1.6         | 21     | 2  | US-08-863-639A-67  |
| 17         | 21    | 1.6         | 21     | 2  | US-08-863-639A-68  |
| 18         | 21    | 1.6         | 21     | 2  | US-08-863-639A-71  |
| 19         | 21    | 1.6         | 21     | 2  | US-08-416-214A-11  |
| 20         | 20    | 1.5         | 20     | 3  | US-09-030-701-65   |
| 21         | 20    | 1.5         | 20     | 4  | US-09-082-649B-57  |
| 22         | 20    | 1.5         | 20     | 4  | US-09-769-482-9    |
| 23         | 20    | 1.5         | 24     | 2  | US-08-570-155-16   |
| 24         | 20    | 1.5         | 24     | 2  | US-08-570-155-17   |
| 25         | 20    | 1.5         | 50     | 3  | US-08-753-247-22   |
| 26         | 20    | 1.5         | 50     | 4  | US-09-930-181-13   |
| 27         | 19    | 1.4         | 28     | 3  | US-09-025-580-6    |

|   |    |     |    |    |   |                   |                    |
|---|----|-----|----|----|---|-------------------|--------------------|
| C | 28 | 1.4 | 19 | 48 | 4 | US-09-438-268-9   | Sequence 9, Appli  |
| C | 29 | 1.4 | 19 | 48 | 4 | US-09-438-268-10  | Sequence 10, Appli |
| C | 30 | 1.4 | 19 | 49 | 1 | US-08-155-171B-27 | Sequence 27, Appl  |
| C | 31 | 1.4 | 19 | 49 | 1 | US-08-155-171B-28 | Sequence 28, Appl  |
| C | 32 | 1.4 | 19 | 49 | 2 | US-08-435-998-27  | Sequence 27, Appl  |
| C | 33 | 1.4 | 19 | 49 | 2 | US-08-435-998-28  | Sequence 28, Appl  |
| C | 34 | 1.4 | 19 | 49 | 4 | US-09-813-781-69  | Sequence 69, Appl  |
| C | 35 | 1.4 | 18 | 49 | 4 | US-09-030-701-65  | Sequence 57, Appl  |
| C | 36 | 1.4 | 18 | 49 | 4 | US-09-082-649B-57 | Sequence 52, Appl  |
| C | 37 | 1.4 | 18 | 21 | 2 | US-08-863-639A-52 | Sequence 55, Appl  |
| C | 38 | 1.4 | 18 | 21 | 2 | US-08-863-639A-55 | Sequence 56, Appl  |
| C | 39 | 1.4 | 18 | 21 | 2 | US-08-863-639A-56 | Sequence 57, Appl  |
| C | 40 | 1.4 | 18 | 21 | 2 | US-08-863-639A-67 | Sequence 67, Appl  |
| C | 41 | 1.4 | 18 | 21 | 2 | US-08-863-639A-68 | Sequence 68, Appl  |
| C | 42 | 1.4 | 18 | 21 | 2 | US-08-863-639A-71 | Sequence 71, Appl  |
| C | 43 | 1.4 | 18 | 21 | 2 | US-08-416-214A-11 | Sequence 11, Appl  |
| C | 44 | 1.4 | 18 | 24 | 2 | US-08-570-155-16  | Sequence 16, Appl  |
| C | 45 | 1.4 | 18 | 24 | 2 | US-08-570-155-17  | Sequence 17, Appl  |

## ALIGNMENTS

RESULT 1  
US-09-475-947A-332  
; Sequence 332, Application US/09475947A  
; Patent No. 6472154  
; GENERAL INFORMATION:  
; APPLICANT: Garner, Harold R.  
; APPLICANT: Wren, Jonathan D.  
; APPLICANT: Minna, John D.  
; TITLE OF INVENTION: Polymorphic Repeats in Human Genes  
; FILE REFERENCE: UTSD0667  
; CURRENT APPLICATION NUMBER: US/09/475,947A  
; CURRENT FILING DATE: 1999-12-31  
; NUMBER OF SEQ ID NOS: 346  
; SOFTWARE: Patentin Ver. 2.1  
; SEQ ID NO 332  
; LENGTH: 30  
; TYPE: DNA  
; ORGANISM: human  
US-09-475-947A-332

Query Match 1.8%; Score 24; DB 4; Length 30;  
Best Local Similarity 100.0%; Pred. No. 0.43;  
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

|    |    |                            |    |
|----|----|----------------------------|----|
| QY | 56 | GGCGGGGGGGGGGGGGGGGGGGGGGA | 79 |
| DB | 7  | GGCGGGGGGGGGGGGGGGGGGGGGGA | 30 |

RESULT 2  
US-09-930-181-14  
; Sequence 14, Application US/09930181  
; Patent No. 6455292  
; GENERAL INFORMATION:  
; APPLICANT: Origene Technologies  
; TITLE OF INVENTION: Full-length Serine Protein Kinase in Brain and Pancreas  
; FILE REFERENCE: 16U 101 V1  
; CURRENT APPLICATION NUMBER: US/09/930,181  
; CURRENT FILING DATE: 2001-08-16  
; NUMBER OF SEQ ID NOS: 18  
; SOFTWARE: Patentin version 3.0  
; SEQ ID NO 14  
; LENGTH: 50  
; TYPE: DNA  
; ORGANISM: Homo sapiens  
US-09-930-181-14

Query Match 1.8%; Score 24; DB 4; Length 50;  
Best Local Similarity 100.0%; Pred. No. 0.41;  
Matches 24; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGGGGGGGGGGGGGGGGGGGG 79  
Db 23 GCGGGGGGGGGGGGGGGGGGG 46

RESULT 3  
US-08-374-144-3  
; Sequence 3, Application US/08374144  
; Patent No. 5629147  
; GENERAL INFORMATION:  
; APPLICANT: Aptogenex, Inc.  
; TITLE OF INVENTION: Enriching and Identifying Fetal Cells  
; TITLE OF INVENTION: Maternal Blood For In Situ Hybridization  
; NUMBER OF SEQUENCES: 21  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Elman Wilf & Fried  
; STREET: 20 West Third Street, P.O. Box 703  
; CITY: Media  
; STATE: PA  
; COUNTRY: USA  
; ZIP: 19063-8969  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: 3.5 inch 720K diskette  
; COMPUTER: IBM PC compatible  
; OPERATING SYSTEM: PC-DOS/MS-DOS  
; SOFTWARE: WordPerfect 5.1  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/374,144  
; FILING DATE:  
; CLASSIFICATION: 435  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Gerry J. Elman  
; REGISTRATION NUMBER: 24,404  
; REFERENCE/DOCKET NUMBER: M19-085  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: 610-892-9580  
; TELEFAX: 610-892-9577  
; INFORMATION FOR SEQ ID NO: 3:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 25 base pairs  
; TYPE: nucleic acid  
; STRANDEDNESS: single  
; TOPOLOGY: linear  
; MOLECULE TYPE: DNA (genomic)  
; HYPOTHETICAL: NO  
; ANTI-SENSE: NO  
US-08-374-144-3

Query Match 1.7%; Score 23; DB 1; Length 25;  
Best Local Similarity 100.0%; Pred. No. 1.2;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGGGGGGGGGGGGGGGGGGGG 78  
Db 2 GCGGGGGGGGGGGGGGGGGGG 24

RESULT 4  
US-08-775-164-3  
; Sequence 3, Application US/08775164  
; Patent No. 576843  
; GENERAL INFORMATION:  
; APPLICANT: Aptogenex, Inc.  
; TITLE OF INVENTION: Enriching and Identifying Fetal Cells  
; NUMBER OF SEQUENCES: 21  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Elman & Associates  
; STREET: 20 West Third Street, P.O. Box 1969  
; CITY: Media  
; STATE: PA  
; COUNTRY: USA  
; ZIP: 19063-8969

Query Match 1.7%; Score 23; DB 1; Length 25;  
Best Local Similarity 100.0%; Pred. No. 1.2;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

COMPUTER READABLE FORM:  
; MEDIUM TYPE: 3.5 inch 720K diskette  
; COMPUTER: IBM PC compatible  
; OPERATING SYSTEM: PC-DOS/MS-DOS  
; SOFTWARE: WordPerfect 5.1  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/775,164  
; FILING DATE:  
; CLASSIFICATION: 530  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Gerry J. Elman  
; REGISTRATION NUMBER: 24,404  
; REFERENCE/DOCKET NUMBER: M19-103  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: 610-892-9580  
; TELEFAX: 610-892-9577  
; INFORMATION FOR SEQ ID NO: 3:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 25 base pairs  
; TYPE: nucleic acid  
; STRANDEDNESS: single  
; TOPOLOGY: linear  
; MOLECULE TYPE: DNA (genomic)  
; HYPOTHETICAL: NO  
; ANTI-SENSE: NO  
US-08-775-164-3

Query Match 1.7%; Score 23; DB 1; Length 25;  
Best Local Similarity 100.0%; Pred. No. 1.2;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGGGGGGGGGGGGGGGGGGGG 78  
Db 2 GCGGGGGGGGGGGGGGGGGGG 24

RESULT 5  
US-08-775-609-3  
; Sequence 3, Application US/08775609  
; Patent No. 5858649  
; GENERAL INFORMATION:  
; APPLICANT: Aptogenex, Inc.  
; TITLE OF INVENTION: Enriching and Identifying Fetal Cells  
; NUMBER OF SEQUENCES: 21  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Elman & Associates  
; STREET: 20 West Third Street, P.O. Box 1969  
; CITY: Media  
; STATE: PA  
; COUNTRY: USA  
; ZIP: 19063-8969  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: 3.5 inch 720K diskette  
; COMPUTER: IBM PC compatible  
; OPERATING SYSTEM: PC-DOS/MS-DOS  
; SOFTWARE: WordPerfect 5.1  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/775,609  
; FILING DATE:  
; CLASSIFICATION: 435  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Gerry J. Elman  
; REGISTRATION NUMBER: 24,404  
; REFERENCE/DOCKET NUMBER: M19-103  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: 610-892-9580  
; TELEFAX: 610-892-9577  
; INFORMATION FOR SEQ ID NO: 3:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 25 base pairs  
; TYPE: nucleic acid  
; STRANDEDNESS: single  
; TOPOLOGY: linear

```
; MOLECULE TYPE: DNA (genomic)
; HYPOTHEICAL: NO
; ANTI-SENSE: NO
US-08-775-609-3
Query Match 1.7%; Score 23; DB 2; Length 25;
Best Local Similarity 100.0%; Pred. No. 1.2;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGCGG 78
DB 2 GCGCGCGCGCGCGCGCGCGCGG 24

RESULT 6
US-08-775-607-3
; Sequence 3, Application US/08775607
; Patent No. 5861253
; GENERAL INFORMATION:
; APPLICANT: Arogenex, Inc.
; TITLE OF INVENTION: Enriching and Identifying Fetal Cells
; NUMBER OF SEQUENCES: 21
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Elman & Associates
; STREET: 20 West Third Street, P.O. Box 1969
; CITY: Media
; STATE: PA
; COUNTRY: USA
; ZIP: 19063-8969
; COMPUTER READABLE FORM:
; MEDIUM TYPE: 3.5 inch 720K diskette
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: WordPerfect 5.1
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/775,607
; FILING DATE:
; CLASSIFICATION: 435
; ATTORNEY/AGENT INFORMATION:
; NAME: Gerry J. Elman
; REGISTRATION NUMBER: 24,404
; REFERENCE/DOCKET NUMBER: M19-103
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 610-892-9580
; TELEFAX: 610-892-9577
; INFORMATION FOR SEQ ID NO: 3:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 25 base pairs
; TYPE: nucleic acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: DNA (genomic)
; HYPOTHETICAL: NO
; ANTI-SENSE: NO
US-08-775-607-3
Query Match 1.7%; Score 23; DB 2; Length 25;
Best Local Similarity 100.0%; Pred. No. 1.2;
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGCGG 78
DB 2 GCGCGCGCGCGCGCGCGCGCGG 24

RESULT 7
PCT-US93-06828-3
; Sequence 3, Application PC/TUS9306828
; GENERAL INFORMATION:
; APPLICANT: Asgari, Morteza
; APPLICANT: Bresser, Joel
; APPLICANT: Cabbage, Michael L
; APPLICANT: Frashad, Nagindra
; TITLE OF INVENTION: Enriching and Identifying Fetal Cells In Maternal Blood For
; NUMBER OF SEQUENCES: 21
; CORRESPONDENCE ADDRESS:
; ADDRESSEE:
; STREET:
; CITY:
; STATE:
; COUNTRY:
; ZIP:
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent In Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/068,747
; FILING DATE: 28-MAY-1993
; CLASSIFICATION: 435
```

ATTORNEY/AGENT INFORMATION:  
NAME: Granahan, Patricia  
REGISTRATION NUMBER: 32,227  
REFERENCE/DOCKET NUMBER: MIT-6141  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: 617-861-6240  
TELEFAX: 617-861-9540  
INFORMATION FOR SEQ ID NO: 2:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 30 base pairs  
TYPE: nucleic acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
MOLECULE TYPE: other nucleic acid  
DESCRIPTION: /desc = "Synthetic"  
US-08-068-747-2

Query Match 1.7%; Score 23; DB 1; Length 30;  
Best Local Similarity 100.0%; Pred. No. 1.1;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
DB 29 GCGCGCGCGCGCGCGCGCGCGG 7

RESULT 9  
US-08-068-747-7  
Sequence 7, Application US/08068747  
Patent No. 5695933  
GENERAL INFORMATION:  
APPLICANT: Schalling, Martin  
APPLICANT: Hudson, Thomas J.  
APPLICANT: Housman, David E.  
TITLE OF INVENTION: Direct Determination of Expanded  
TITLE OF INVENTION: Nucleotide Repeats in the Human Genome  
NUMBER OF SEQUENCES: 11  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: Hamilton, Brook, Smith & Reynolds, P.C.  
STREET: Two Militia Drive  
CITY: Lexington  
STATE: Massachusetts  
COUNTRY: USA  
ZIP: 02173  
COMPUTER READABLE FORM:  
MEDIUM TYPE: Floppy disk  
COMPUTER: IBM PC Compatible  
OPERATING SYSTEM: PC-DOS/MS-DOS  
SOFTWARE: Patent in Release #1.0, Version #1.30  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/08/068,747  
FILING DATE: 28-MAY-1993  
CLASSIFICATION: 435  
ATTORNEY/AGENT INFORMATION:  
NAME: Granahan, Patricia  
REGISTRATION NUMBER: 32,227  
REFERENCE/DOCKET NUMBER: MIT-6141  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: 617-861-6240  
TELEFAX: 617-861-9540  
INFORMATION FOR SEQ ID NO: 7:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 33 base pairs  
TYPE: nucleic acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
MOLECULE TYPE: other nucleic acid  
DESCRIPTION: /desc = "Synthetic"  
US-08-068-747-7

Query Match 1.7%; Score 23; DB 1; Length 33;  
Best Local Similarity 100.0%; Pred. No. 1.1;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
DB 2 GCGCGCGCGCGCGCGCGCGCGG 24

RESULT 10  
US-09-497-933A-22/c  
Sequence 22, Application US/09497933A  
Patent No. 6329147  
GENERAL INFORMATION:  
APPLICANT: Wagner, Robert Jr. E.  
TITLE OF INVENTION: METHODS FOR DETECTION OF A TRIPLET REPEAT BLOCK AND A  
TITLE OF INVENTION: FUNCTIONAL MISMATCH BINDING PROTEIN IN A BIOLOGICAL  
TITLE OF INVENTION: FLUID SAMPLE  
FILE REFERENCE: 9408-044  
CURRENT APPLICATION NUMBER: US/09/497,933A  
CURRENT FILING DATE: 2000-02-04  
NUMBER OF SEQ ID NOS: 25  
SOFTWARE: Patent in Ver. 2.1  
SEQ ID NO 22  
LENGTH: 45  
TYPE: DNA  
ORGANISM: Artificial Sequence  
FEATURE:  
OTHER INFORMATION: Description of Artificial Sequence: Probe  
NAME/KEY: modified\_base  
LOCATION: (1)..(9)  
OTHER INFORMATION: n = a, c, g or t  
NAME/KEY: modified\_base  
LOCATION: (37)..(45)  
OTHER INFORMATION: n = a, c, g or t  
US-09-497-933A-22

Query Match 1.7%; Score 23; DB 4; Length 45;  
Best Local Similarity 100.0%; Pred. No. 1.1;  
Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGG 78  
DB 35 GCGCGCGCGCGCGCGCGCGCGG 13

RESULT 11  
US-09-497-933A-19  
Sequence 19, Application US/09497933A  
Patent No. 6329147  
GENERAL INFORMATION:  
APPLICANT: Wagner, Robert Jr. E.  
TITLE OF INVENTION: METHODS FOR DETECTION OF A TRIPLET REPEAT BLOCK AND A  
TITLE OF INVENTION: FUNCTIONAL MISMATCH BINDING PROTEIN IN A BIOLOGICAL  
TITLE OF INVENTION: FLUID SAMPLE  
FILE REFERENCE: 9408-044  
CURRENT APPLICATION NUMBER: US/09/497,933A  
CURRENT FILING DATE: 2000-02-04  
NUMBER OF SEQ ID NOS: 25  
SOFTWARE: Patent in Ver. 2.1  
SEQ ID NO 19  
LENGTH: 48  
TYPE: DNA  
ORGANISM: Artificial Sequence  
FEATURE:  
OTHER INFORMATION: Description of Artificial Sequence: Probe  
NAME/KEY: modified\_base  
LOCATION: (1)..(9)  
OTHER INFORMATION: n = a, c, g or t  
NAME/KEY: modified\_base  
LOCATION: (40)..(48)  
OTHER INFORMATION: n = a, c, g or t  
US-09-497-933A-19

Query Match 1.7%; Score 23; DB 4; Length 48;  
Best Local Similarity 100.0%; Pred. No. 1.1;

Matches 23; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 GCGCGCGCGCGCGCGCGCGCGCG 78  
 Db 11 GCGCGCGCGCGCGCGCGCGCGCG 33

RESULT 12  
 US-09-769-482-8/c  
 ; Sequence 8, Application US/09769482  
 ; Patent No. 6566130  
 ; GENERAL INFORMATION:  
 ; APPLICANT: SRIVASTAVA, SHIV  
 ; APPLICANT: MOUL, JUDU W.  
 ; APPLICANT: XU, LINDA L.  
 ; APPLICANT: SEGAWA, TAKEHIKO  
 ; TITLE OF INVENTION: PROSTATE-SPECIFIC ANDROGEN-SIGNALING-ASSOCIATED  
 ; FILE REFERENCE: POYNUCLEOTIDE ARRAY  
 ; CURRENT APPLICATION NUMBER: US/09/769,482  
 ; CURRENT FILING DATE: 2001-01-26  
 ; PRIOR APPLICATION NUMBER: 60/178,772  
 ; PRIOR FILING DATE: 2000-01-26  
 ; PRIOR APPLICATION NUMBER: 60/179,045  
 ; PRIOR FILING DATE: 2000-01-31  
 ; NUMBER OF SEQ ID NOS: 67  
 ; SOFTWARE: PatentIn Ver. 2.1  
 ; SEQ ID NO 8  
 ; LENGTH: 22  
 ; TYPE: DNA  
 ; ORGANISM: Artificial Sequence  
 ; FEATURE:  
 ; OTHER INFORMATION: Description of Artificial Sequence: Primer  
 US-09-769-482-8

Query Match 1.7%; Score 22; DB 4; Length 22;  
 Best Local Similarity 100.0%; Pred. No. 3.2;  
 Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 838 CGAGATCGACCTGCCACCCACC 859  
 Db 22 CGAGATCGACCTGCCACCCACC 1

RESULT 13  
 US-08-863-639A-52/c  
 ; Sequence 52, Application US/08863639A  
 ; Patent No. 5981185  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Matson, Robert S.  
 ; APPLICANT: Coassin, Peter J.  
 ; APPLICANT: Rampal, Jang B.  
 ; APPLICANT: Caskey, C. T.  
 ; TITLE OF INVENTION: OLIGONUCLEOTIDE REPEAT ARRAYS  
 ; NUMBER OF SEQUENCES: 95  
 ; CORRESPONDENCE ADDRESS:  
 ; ADDRESSEE: Sheldon & Mak  
 ; STREET: 225 South Lake Avenue, 9th Floor  
 ; CITY: Pasadena  
 ; STATE: CA  
 ; COUNTRY: USA  
 ; ZIP: 91101  
 ; COMPUTER READABLE FORM:  
 ; MEDIUM TYPE: Diskette, 3.50 inch, 1.44 Mb storage  
 ; COMPUTER: IBM compatible  
 ; OPERATING SYSTEM: Windows 95  
 ; SOFTWARE: Corel WordPerfect 8 version  
 ; CURRENT APPLICATION DATA:  
 ; APPLICATION NUMBER: US/08/863.639A  
 ; FILING DATE: May 28, 1997  
 ; CLASSIFICATION: 435  
 ; ATTORNEY/AGENT INFORMATION:  
 ; NAME: Joseph E. Mueth

REGISTRATION NUMBER: 20,532  
 REFERENCE/DOCKET NUMBER: 11859-1  
 TELECOMMUNICATION INFORMATION:  
 TELEPHONE: (626) 796-4000  
 TELEFAX: (626) 795-6321  
 INFORMATION FOR SEQ ID NO: 52:  
 SEQUENCE CHARACTERISTICS:  
 LENGTH: 21 base pairs  
 TYPE: nucleic acid  
 STRANDEDNESS: single  
 TOPOLOGY: linear  
 MOLECULE TYPE: Other nucleic acid  
 US-08-863-639A-52

Query Match 1.6%; Score 21; DB 2; Length 21;  
 Best Local Similarity 100.0%; Pred. No. 8.8;  
 Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 58 CGCGCGCGCGCGCGCGCGCGCG 78  
 Db 21 CGCGCGCGCGCGCGCGCGCGCG 1

RESULT 14  
 US-08-863-639A-55/c  
 ; Sequence 55, Application US/08863639A  
 ; Patent No. 5981185  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Matson, Robert S.  
 ; APPLICANT: Coassin, Peter J.  
 ; APPLICANT: Rampal, Jang B.  
 ; APPLICANT: Caskey, C. T.  
 ; TITLE OF INVENTION: OLIGONUCLEOTIDE REPEAT ARRAYS  
 ; NUMBER OF SEQUENCES: 95  
 ; CORRESPONDENCE ADDRESS:  
 ; ADDRESSEE: Sheldon & Mak  
 ; STREET: 225 South Lake Avenue, 9th Floor  
 ; CITY: Pasadena  
 ; STATE: CA  
 ; COUNTRY: USA  
 ; ZIP: 91101  
 ; COMPUTER READABLE FORM:  
 ; MEDIUM TYPE: Diskette, 3.50 inch, 1.44 Mb storage  
 ; COMPUTER: IBM compatible  
 ; OPERATING SYSTEM: Windows 95  
 ; SOFTWARE: Corel WordPerfect 8 version  
 ; CURRENT APPLICATION DATA:  
 ; APPLICATION NUMBER: US/08/863.639A  
 ; FILING DATE: May 28, 1997  
 ; CLASSIFICATION: 435  
 ; ATTORNEY/AGENT INFORMATION:  
 ; NAME: Joseph E. Mueth

REGISTRATION NUMBER: 20,532  
 REFERENCE/DOCKET NUMBER: 11859-1  
 TELECOMMUNICATION INFORMATION:  
 TELEPHONE: (626) 796-4000  
 TELEFAX: (626) 795-6321  
 INFORMATION FOR SEQ ID NO: 55:  
 SEQUENCE CHARACTERISTICS:  
 LENGTH: 21 base pairs  
 TYPE: nucleic acid  
 STRANDEDNESS: single  
 TOPOLOGY: linear  
 MOLECULE TYPE: Other nucleic acid  
 US-08-863-639A-55

Query Match 1.6%; Score 21; DB 2; Length 21;  
 Best Local Similarity 100.0%; Pred. No. 8.8;  
 Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 57 GCGCGCGCGCGCGCGCGCGCGCG 77  
 Db 21 GCGCGCGCGCGCGCGCGCGCGCG 1

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RESULT 15
US-08-863-639A-56
; Sequence 56, Application US/08863639A
; Patent No. 5981185
; GENERAL INFORMATION:
; APPLICANT: Matson, Robert S.
; APPLICANT: Coassin, Peter J.
; APPLICANT: Rampal, Jang B.
; APPLICANT: Caskey, C.T.
; TITLE OF INVENTION: OLIGONUCLEOTIDE REPEAT ARRAYS
; NUMBER OF SEQUENCES: 95
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Sheldon & Mak
; STREET: 225 South Lake Avenue, 9th Floor
; CITY: Pasadena
; STATE: CA
; COUNTRY: USA
; ZIP: 91101
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette, 3.50 inch, 1.44 Mb storage
; COMPUTER: IBM compatible
; OPERATING SYSTEM: Windows 95
; SOFTWARE: Corel WordPerfect 8 version
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/863,639A
; FILING DATE: May 28, 1997
; CLASSIFICATION: 435
; ATTORNEY/AGENT INFORMATION:
; NAME: Joseph E. Mueth
; REGISTRATION NUMBER: 20,532
; REFERENCE/DOCKET NUMBER: 11859-1
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (626) 796-4000
; TELEFAX: (626) 795-6321
; INFORMATION FOR SEQ ID NO: 56:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 21 base pairs
; TYPE: nucleic acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: Other nucleic acid
; US-08-863-639A-56

```

```

Query Match      1.6%; Score 21; DB 2; Length 21;
Best Local Similarity 100.0%; Pred. No. 8.8;
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

```

```

Qy      58  CGGCGGCGGCGGCGGCGGCGG 78
Db      1  CGGCGGCGGCGGCGGCGGCGG 21

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Search completed: July 1, 2004, 21:53:07
Job time : 115 secs

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